

UNIVERSIDADE TÉCNICA DE LISBOA INSTITUTO SUPERIOR TÉCNICO



Salt crystallization in plastered or rendered walls

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(MSc)

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Salt crystallization in plastered or rendered walls

Abstract

This thesis was aimed at understanding the behaviour of plasters and renders on salt-loaded walls. The current state-of-the-art and state-of-the-practice were accessed focusing particularly on old plastered/rendered buildings and their conservation practice in Portugal. Afterwards, experimental work was carried out aiming at answering the identified questions. Two laboratory techniques, for relative humidity control with salt solutions and for salt content determination by hygroscopic moisture content measurements, were investigated. Drying of salt-loaded materials was studied by means of drying experiments monitored using a magnetic resonance imaging (MRI) technique. The behaviour of plasters and renders in relation to salt crystallization was then accessed by means of crystallization tests and MRI-monitored drying tests. Most drying and crystallization tests were carried out on specimens composed by a plaster or render applied on a given substrate. Finally, the study of five old buildings in Portugal provided an insight into practice-related salt decay features.

On the basis of this research, guidelines are proposed to select plasters and renders for saltloaded walls. Conclusions were also achieved on: (i) possibilities and limitations of the test methods, particularly salt crystallization tests, and diagnostic methodology used; (ii) salt decay processes, namely, influence of soluble salts on drying, mechanisms of salt-induced dampness and salt distribution in masonry; (iii) reasons for sodium chloride being typically much less damaging than sodium sulfate in laboratory tests; (iv) influence of factors such as the type of salt, kind of substrate material or presence of a paint layer on the behaviour of plasters and renders; (v) factors that can account for a worsening of salt damage after restoration interventions; (vi) field or application conditions that favour salt damage.

Key-words: soluble salt crystallization, salt decay, plasters and renders, old buildings, salt-damp, efflorescence

Cristalização de sais solúveis em paredes rebocadas

Resumo

A presente tese visa compreender o comportamento de rebocos aplicados em paredes contaminadas com sais solúveis. O estado-da-arte e o estado-da-prática foram avaliados focando, em particular, o caso dos edifícios antigos e a situação da prática em Portugal. O trabalho experimental seguidamente realizado visou dar resposta às principais questões identificadas. Foram avaliadas duas técnicas laboratoriais de interesse geral, para controlo da humidade relativa por meio de soluções salinas e para avaliação do teor de sal com base na medição do teor de humidade higroscópica. A secagem de materiais contaminados com sais foi investigada através de ensaios de secagem monitorizados por ressonância magnética nuclear. O comportamento de rebocos relativamente à cristalização de sais for estudado por meio de ensaios de cristalização e de ensaios de secagem monitorizados por ressonância magnética magnética nuclear. A maioria dos ensaios, quer de cristalização, quer de secagem, foram realizados em provetes compostos por um reboco aplicado num determinado suporte. Finalmente, foram estudados cinco edifícios antigos portugueses, o que possibilitou compreender causas e tipos de degradação que frequentemente ocorrem na prática.

A tese permitiu estabelecer orientações para a selecção de rebocos para paredes contaminadas com sais solúveis. Permitiu ainda retirar conclusões sobre: (i) possibilidades e limitações dos métodos de ensaio, em particular de cristalização de sais, e metodologia de diagnóstico utilizados; (ii) processos de degradação, nomeadamente, influência dos sais solúveis na secagem de materiais porosos, problemas de humidade devidos a sais solúveis e distribuição de sal em alvenarias; (iii) razões para o cloreto de sódio provocar tipicamente maior degradação do que o sulfato de sódio em ensaios laboratoriais de cristalização; (iv) influência no comportamento dos rebocos de factores como o tipo de sal, natureza dos materiais do suporte ou presença de pintura; (v) factores que podem originar um agravamento dos danos por sais solúveis após intervenções de conservação; (vi) condições de obra que favorecem a degradação por sais solúveis.

Palavras-chave: cristalização de sais solúveis, degradação por sais solúveis, rebocos, edifícios antigos, humidade em edifícios, eflorescências

List of contents

Chapter 1 – Introduction

1.1	Motivation and scope	.1
1.2	Aims and objectives	.3
1.3	Outline of the thesis	.4
1.4	Delimitations of scope and key-assumptions	.5

Chapter 2 – Research and practice: current situation

2.1	Introduction	7
2.2	State of the art	8
	2.2.1 - Salt decay pathology	8
	2.2.2 - Signs	9
	2.2.3 - Moisture	11
	2.2.4 - Soluble salts	12
	2.2.5 - Damage mechanisms	16
	2.2.6 - Influencing factors	19
	2.2.7 - Working principles of plasters and renders	22
	2.2.8 - Salt crystallization tests	24
2.3	State of the practice	24
	2.3.1 - Introduction	24
	2.3.2 - General approach to salt decay problems	25
	2.3.3 - Main types of plasters and renders used on salt-loaded walls	26

Chapter 3 – General laboratory techniques

3.1 Introduction	31
3.2 Fundamentals	32
3.2.1 - Relative equilibrium humidity of aqueous solutions	32
3.2.2 - Control of the relative humidity by means of aqueous solutions	33
3.2.3 - Hygroscopic behaviour of soluble salts	33
3.3 Use of aqueous solutions for control of the relative humidity	34
3.3.1 - Introduction	34

3.3.2 - Materials and methods	35
3.3.3 - Results	35
3.3.4 - Discussion	36
3.4 Fundamentals, scope and accuracy of the HMC method	
3.4.1 - Introduction	38
3.4.2 - Systematization of the HMC method	40
3.4.2.1 Measurement of the HMC	40
3.4.2.2 Determination of the salt content	40
3.4.3 - Discussion	42
3.4.3.1 Accuracy of HMC measurements	42
3.4.3.2 Mass stabilization	44
3.4.3.3 Scope of the HMC method	45
3.5 HMC experiments on salt-loaded materials	46
3.5.1 - Introduction	46
3.5.2 - Materials and methods	48
3.5.3 - Results	50
3.5.3.1 - General	50
3.5.3.2 - Mass stabilization	51
3.5.3.3 - Pure salts	51
3.5.3.4 - Base-materials (mortar and brick)	53
3.5.3.5 - Sodium chloride or sodium sulphate loaded materials: HMC at 80% RH	53
3.5.3.6 - Sodium chloride loaded materials	54
3.5.3.7 - Sodium sulphate loaded materials	54
3.5.3.8 - Mortar loaded with other soluble salts	56
3.5.3.9 - Salt mixtures	57
3.5.3.10 - Error in salt content estimates	59
3.5.4 - Discussion	60
3.5.4.1 - Influence of the base-materials and their state of cohesion	60
3.5.4.2 - Mass stabilization	60
3.5.4.3 - Correlation HMC/salt content	61
3.5.4.4 - Accuracy	61
3.5.4.5 - Sodium sulphate	62
3.5.4.6 - Salt mixtures	63
3.6 Conclusions	64

Chapter 4 – Drying of salt-contaminated materials

4.1 Introduction	69
4.2 Drying of porous materials	71
4.2.1 - Porosity and hygroscopicity	71
4.2.2 - Liquid water transport	72
4.2.3 - Water vapour transport	77
4.2.4 - Drying of porous materials	78

4.3 Materials and methods	81
4.3.1 - Materials	81
4.3.2 - Drying experiments	83
4.3.3 - Salt distribution	83
4.4 Results	84
4.4.1 - Drying of water-filled specimens	84
4.4.2 - Drying in the presence of NaCl	86
4.4.3 - Overall drying curves	87
4.4.4 - Evaporation from free liquid surfaces	87
4.4.5 - Salt deposition	88
4.5 Discussion	89
4.5.1 - Salt-induced dampness	89
4.5.2 - Salt distribution	91
4.5.3 - Other soluble salts	93

Chapter 5 – Behaviour of plasters and renders

5.1	Introduction	95
5.2	Salt crystallization tests	98
	5.2.1 - Introduction	98
	5.2.2 - Test variables	99
	5.2.3 - Type of specimens	100
	5.2.4 - Contamination conditions	101
	5.2.5 - Drying procedure	102
	5.2.6 - Salt solutions	103
	5.2.7 - Damage assessment	104
5.3	Sorption of porous building materials	104
5.4	Material characterization	106
5.5	Crystallization tests on plaster/substrate specimens (set 1)	109
	5.5.1 - Introduction	109
	5.5.2 - Experimental conditions	109
	5.5.2.1 - Definition of the main testing protocol	109
	5.5.2.2 - Materials and methods	111
	5.5.3 - Results	113
	5.5.3.1 - Salt crystallization tests	113
	5.5.3.2 - Tests with pure water	118
	5.5.3.3 - Salt introduced in the specimens	118
	5.5.3.4 - Drying curves	119
	5.5.3.5 - Summary of results	121
	5.5.4 - Discussion on set 1	122
	5.5.4.1 - Main behaviour features	122
	5.5.4.2 - Influence of the type of salt	124
	5.5.4.3 - Influence of the paint	126

5.5.4.4 - Influence of the substrate material	
5.5.4.5 - Test method	
5.6 Drying of a salt-accumulating plaster (set 2)	
5.6.1 - Introduction	
5.6.2 - Materials and methods	
5.6.3 - Results	131
5.6.4 - Discussion on set 2	
5.6.4.1 - Totally saturated specimens	
5.6.4.2 - Partially saturated specimens	
5.7 Influence of a paint on drying (set 3)	140
5.7.1 - Introduction	140
5.7.2 - Materials and methods	140
5.7.3 - Results	140
5.7.4 - Discussion on set 3	142
5.8 Drying of hydrophobic agent containing plaster (set 4)	
5.8.1 - Introduction	
5.8.2 - Materials and methods	
5.8.3 - Results	144
5.8.4 - Discussion on set 4	
5.9 Crystallization tests on painted stone (set 5)	147
5.9.1 - Introduction	147
5.9.2 - Materials and methods	147
5.9.3 - Results	
5.9.4 - Discussion on set 5	
5.10 Discussion	151
5.10.1 - Salt distribution	151
5.10.2 - Plasters and renders	
5.10.3 - Sodium chloride and sodium sulphate	
5.10.4 - Development of an accelerated-ageing test	157
5.10.5 - Restoration interventions	
5.10.6 - Adequability of plasters and renders	

Chapter 6 – Case studies

6.1	Introduction	161
6.2	Description of the buildings and main decay patterns	163
	6.2.1 - Salvas Chapel	163
	6.2.2 - Alhos-Vedros tide mill	165
	6.2.3 - Despacho House	167
	6.2.4 - Cloister of Sta Clara-a-Nova Monastery	168
	6.2.5 - S. Sebastião Church	171
6.3	Methods	172

6.4 Results and discussion	173
6.4.1 - General	173
6.4.2 - Salvas Chapel	174
6.4.3 - Alhos-Vedros tide mill	175
6.4.4 - Despacho House	177
6.4.5 - Cloister of Sta Clara-a-Nova Monastery	179
6.4.6 - S. Sebastião Church	182
6.5 Research on experimental test panels	183
6.5.1 - Materials and methods	183
6.5.2 - Execution of the test panels	185
6.5.3 - Evaluation of the systems after three years of natural exposure	187
6.6 Discussion	189
6.6.1 - Main causes of decay	189
6.6.2 - Diagnosis	191
6.6.3 - Test panels	191

Chapter 7 – Summary, conclusions and future perspectives

7.1	Summary	193
7.2	Conclusions	195
	7.2.1 - Test methods	195
	7.2.2 - Salt decay processes	197
	7.2.3 - Sodium chloride versus sodium sulphate	198
	7.2.4 - Factors that influence salt damage features	199
	7.2.5 - Factors that can contribute for an aggravation of salt damage	200
	7.2.6 - Other field conditions that favour salt damage	200
	7.2.7 - Diagnostic methodology	201
	7.2.8 - Application technique of industrial plasters and renders	201
	7.2.9 - Selecting plasters and renders for moist salt-loaded walls	201
7.3	Future perspectives	203
Ref	ferences	207

Annex I – Magnetic resonance imaging

I.1	Introduction	AI.1
1.2	One-dimensional technique	AI.3
1.3	Two-dimensional technique	AI.5

Annex II – Making of the test specimens for set 1 tests

Annex III – Damage assessment and diagnosis form

List of figures

Chapter 1 – Introduction

Fia.	1.1 -	Two typica	l salt-damage	patterns	1
i ig.	1.1 -	i wo typico	a san-uamage	pationis	 •

Chapter 2 – Research and practice: current situation

Fig. 2.1 – Salt decay pathology	8
Fig. 2.2 – Some common salt decay patterns of plasters and renders	10
Fig. 2.3 – Alveolar decay	11
Fig. 2.4 – Masonry disintegration	11
Fig. 2.6 – Solubility diagram of sodium chloride	15
Fig. 2.7 – RH/T phase diagram of sodium chloride	15
Fig. 2.8 – Solubility diagram of sodium sulphate	16
Fig. 2.9 – RH/T phase diagram of sodium sulphate	16
Fig. 2.10 – Representation of the four main working principles of plasters and renders	23

Chapter 3 – General laboratory techniques

33
36
36
36
36
41
41
42
43
44
50
51
52
52

Fig. 3.15 – HMC of the base materials at 95% RH and 20°C	53
Fig. 3.16 – HMC at 80% RH of mortar and brick samples loaded with NaCl or Na_2SO_4	54
Fig. 3.17 – HMC at 95% RH of NaCl-loaded brick or mortar samples	54
Fig. 3.18 – HMC at 95% RH of Na ₂ SO ₄ -loaded brick or mortar samples	55
Fig. 3.19 – Evolution over time of the HMC of mortar samples with up to 100% Na_2SO_4	
Fig. 3.20 – HMC at 95% RH of mortar samples loaded with each of six distinct salts	57
Fig. 3.21 – HMC of mortar loaded with salt mixtures	57
Fig. 3.22 – Actual HMC for the salt mixtures and theoretical HMC	

Chapter 4 – Drying of salt-contaminated materials

Fig. 4.1 – Capillary pressure in a cylindrical capillary	72
Fig. 4.2 – Porous material in free absorption	75
Fig. 4.3 – Typical water absorption curves for porous building materials	75
Fig. 4.4 – Moisture transport at the pore scale during wetting of a porous material	79
Fig. 4.5 – Drying of a porous material at a macroscopic scale	79
Fig. 4.6 – Typical drying curve of a porous building material	80
Fig. 4.7 – Pictures of one specimen	81
Fig. 4.8 – Pore size distribution (MIP)	82
Fig. 4.9 – Drying with pure water: MRI snapshots	85
Fig. 4.10 – Drying with pure water: 1D moisture profiles at cross-sections CS1 and CS2	85
Fig. 4.11 – MRI snapshots of specimen DL1 (side-view) during drying with: (a) pure water;	
(b) 3m NaCl solution.	86
Fig. 4.12 – MRI snapshots of specimen ML1 (side-view) during drying with: (a) pure water;	
(b) 3m NaCl solution	86
Fig. 4.13 – Overall drying curves	87
Fig. 4.14 – Evaporation from free surfaces of pure water and saturated NaCl solution	87
Fig. 4.15 – Final salt distribution in specimen DL1	88
Fig. 4.16 – Final salt distribution in specimen ML1	88

Chapter 5 – Behaviour of plasters and renders

Fig. 5.1 – Crystallization test on brick treated with hydrophobic surface treatment on all faces	
except the bottom (specimen I) and non-treated brick (specimen II)	96
Fig. 5.2 – Test panels on the interior of S. Sebastião church	97
Fig. 5.3 – Preliminary crystallization tests by wet-dry cycles	103
Fig. 5.4 – Water characteristic curve of a porous building material	105
Fig. 5.5 – Capillary absorption tests - WAC in kg/(m ² .h ^{1/2})	107
Fig. 5.6 – Drying curves	107
Fig. 5.7 – Pore size distribution (MIP)	108
Fig. 5.8 – Water vapour transport.	108
Fig. 5.9 – Preliminary crystallization tests by continuous immersion	110
Fig. 5.10 – Specimens (side view) used in set 1 tests	111
Fig. 5.11 – HMC profiles of some non-painted specimens tested with pure water	118

Fin 5.40 MED as heidlich in a sum a	440
Fig. 5.12 – MEP on brick drying curves	119
Fig. 5.13 – Parlumière on brick drying curves	119
Fig. 5.14 – LNEC on brick drying curves	120
Fig. 5.15 – LNEC on lime mortar drying curves	120
Fig. 5.16 – Drying curves of the three LNEC on lime mortar specimens with sodium sulphate	120
Fig. 5.17 – Correlation HMC / ion content	130
Fig. 5.18 – Results of the MRI-monitored drying experiments on fully saturated specimens	132
Fig. 5.19 – Drying curves: total amount of water in the saturated specimens	132
Fig. 5.20 - Results of the MRI-monitored drying experiments on partially saturated specimens	133
Fig. 5.21 – Drying curves: total amount of water in the partially saturated specimens	133
Fig. 5.22 - Difference between evaporation and liquid fluxes in the partially saturated specimens	133
Fig. 5.23 – Liquid fluxes in the partially saturated specimens	134
Fig. 5.24 – Evaporation fluxes in the partially saturated specimens	134
Fig. 5.25 – Chloride distribution in MEP 2	134
Fig. 5.26 – Sulphate distribution in MEP 1	134
Fig. 5.27 – Representation of liquid migration in MEP plaster	137
Fig. 5.28 – Representation of the specimen assemblage (side-view) and cross-sections	140
Fig. 5.29 – MRI snapshots of specimens D2 and D1P during drying with pure water	141
Fig. 5.30 – Moisture content profiles at cross-sections NP (above) and P (below)	141
Fig. 5.31 – Drying curves	141
Fig. 5.32 – Drying with pure water: MRI snapshots of specimens DL1 and DH3 on side-view	144
Fig. 5.33 – Drying with pure water: 1D moisture profiles at cross-sections CS1 and CS2 of	
specimens DH3 and DL1	144
Fig. 5.34 – Normalized drving curves	145
Fig. 5.35 – Gravimetric drving experiments on water-filled specimens of non-additivated	
plaster L or additivated plaster H	
Fig. 5.36 – Damage on the acrylic paint at the end of the first crystallization experiment.	
Fig. 5.37 – Damage on the non-painted stone specimens at the end of the first crystallization	
experiment	149
Fig. 5.38 – Damage to the acrylic paint during the second crystallization experiment	140
Fig. 5.39 – Paint damage arising without the presence of efflorescence	156
Fig. 5.40 $-$ Paint damage by salt efflorescence	156
T 19. 0. TO = 1 and damage by sail emolescence	

Chapter 6 – Case studies

Fig. 6.1 – Salvas Chapel, SW (front) façade	163
Fig. 6.2 – Salvas Chapel, NE (rear) façade	164
Fig. 6.3 – Salvas Chapel, SE façade	164
Fig. 6.4 – Salvas Chapel, NW façade	164
Fig. 6.5 – Salvas Chapel, damage spots on the SE façade	164
Fig. 6.6 – Dona Bataça (water fountain plus reservoir)	164
Fig. 6.7 – Tide-mill (N façade)	165
Fig. 6.8 – Tide-mill, damage at the ground floor, northern wall	166
Fig. 6.9 – Tide-mill, damage on the southern wall	166

Fig. 6.10 - Tide-mill: moisture stains, probably due to dew point condensation, on the ston	e
pavement of the ground floor	166
Fig. 6.11 – Pereira Misericórdia West façade	167
Fig. 6.12 – Extensive damage at the interior of Despacho House	168
Fig. 6.13 – Sta. Clara-a-Nova cloister gallery	169
Fig. 6.14 – Sta. Clara: damage appears either close to the ground (NE wall, on the left)	
or higher on the walls, next to stone elements (NW wall, on the right)	169
Fig. 6.15 - Sta. Clara: details of lower (on the left) and upper (on the centre) damage spots	s169
Fig. 6.16 – Sta. Clara: sequence of a possible damage mechanism	170
Fig. 6.17 – Sta. Clara: sporadic cracks are observed	170
Fig. 6.18 – Sta. Clara: in some cases, damage starts by the bulging and further detachment	nt
of a thin surface layer of the render, accompanied by peeling of the paint, without	it the
presence of efflorescence	170
Fig. 6.19 - Sta. Clara: moisture spots on the stone (left), moisture spots (centre) and biolog	gical
growth (right) on the render	170
Fig. 6.20 - Sta. Clara: on the same day in February 2004 probable dew point condensation	า
events were also observed on the stone floor (left and centre). Accumulation	
of water in the cloister garden (right) had been observed during a previous	
inspection, in January 2004	171
Fig. 6.21 – São Sebastião Church: NE (front) façade in 1999	171
Fig. 6.22 – São Sebastião: masonry damage	172
Fig. 6.23 – Assessment of the possible water loss from the drilling technique	173
Fig. 6.24 – Salvas rear façade: drilling location (left) and MC/HMC profiles (centre and right	t)174
Fig. 6.25 – Tide mill NW wall: drilling location (left) and MC/HMC profiles (centre and right)	176
Fig. 6.26 – Despacho internal wall: drilling location (left) and MC/HMC profiles (centre and	right)178
Fig. 6.27 - Sta Clara NW wall: drilling location (left) and MC/HMC profiles (centre and right	t)179
Fig. 6.28 – Sta Clara NE wall: drilling location (left) and MC/HMC profiles (centre and right)180
Fig. 6.29 - S. Sebastião: drilling location (left) and MC/HMC profiles (centre and right)	182
Fig. 6.30 – Test panels: traditional renders A and C and industrial renders D, E and F	183
Fig. 6.31 - Capillary water absorption of the three industrial systems and traditional mortan	A184
Fig. 6.32 – Before making the panels, the substrate was prepared	185
Fig. 6.33 – Test panel execution	186
Fig. 6.34 – Cracking of the rendering systems three years after their application	
Fig. 6.35 – Comparison between the salt content of the masonry materials	

Annex I – Magnetic resonance imaging

Fig. I.1 – NMR set-up for 1D measurements at TUE	AI.3
Fig. I.2 – Test-specimen sealed with Teflon tape	AI.4
Fig. I.3 – NMR coil, with specimen inside the central sample-holder, and air-flow tube	AI.4
Fig. I.4 – Moisture distribution profiles obtained during one drying experiment (above) and	
representation of the measured specimen (bellow)	AI.4

Fig. I.5 – Correlation moisture content / NMR signal for MEP plaster and brick	. Al.5
Fig. I.6 – NMR set-up for 2D measurements at TUE	. AI.5
Fig. I.7 – Test-specimen made of plaster on mortar/stone substrate	. AI.6
Fig. I.8 – Sample-holder with specimen inside	. AI.6
Fig. I.9 – NMR coil with sample-holder connected to air-flow	. AI.6
Fig. I.10 – Representation of 2D measurement on a sample	. AI.7
Fig. I.11 – Example of a 2D NMR image	. AI.7
Fig. I.12 – Correlation moisture content / NMR signal for plaster on mortar + stone specimens	. AI.8
Fig. I.13 – Colour scale for the 2D NMR measurements	. AI.8

List of tables

Chapter 2 – Research and practice: current situation

Table 2.1 – Requirements for moisture transport properties of CEN renovation mortars	
and WTA sanierputz	28

Chapter 3 – General laboratory techniques

Table 3.1 – HMC experiments	48
Table 3.2 – Relative equilibrium humidity of the salts	49
Table 3.3 – Composition of the salt mixtures	49
Table 3.4 – Actual RH during the second set: at the end of the tests on loaded brick or mortar	
(29 days) and on pure salt samples (64 days)	53
Table 3.5 - HMC at 80%RH and 20°C: actual RH and correlation equations for NaCI-loaded	
brick or mortar samples	54
Table 3.6 - HMC at 95%RH and 20°C: actual RH and correlation equations for NaCI-loaded	
brick or mortar samples	54
Table 3.7 – HMC at 95%RH and 20°C: actual RH and correlation equations for mortar or	
brick samples with Na_2SO_4 -content below 2%	55
Table 3.8 – HMC at 95%RH and 20°C: actual RH and correlation equations for mortar samples	
loaded with six distinct salts	57
Table 3.9 – HMC at 95%RH and 20°C: correlation equations for mortar samples loaded	
with four salt mixtures	57
Table 3.10 – Agreement of the actual HMC of the mixture with the value obtained by weighting	
the HMC of the individual salts	58
Table 3.11 – Significance of the maximum HMC variation among the salt mixtures	59
Table 3.12 – Error in salt content estimates	59

Chapter 4 – Drying of salt-contaminated materials

Table 4.1 – Composition	, porosity and capillar	y coefficient of the materials	82
-------------------------	-------------------------	--------------------------------	----

Chapter 5 – Behaviour of plasters and renders

Table 5.1 – Materials used in the specimens	106
Table 5.2 – Environmental conditions used in the preliminary crystallization tests	110
Table 5.3 – Set 1 crystallization tests	112
Table 5.4 – Crystallization test on MEP-SP / brick specimens	114
Table 5.5 – Crystallization test on Parlumière / brick specimens	115
Table 5.6 – Crystallization test on LNEC / brick specimens	116
Table 5.7 – Crystallization test on LNEC / lime-mortar specimens	117
Table 5.8 – Salt introduced in the specimens	118
Table 5.9 – NaCI crystallization tests (set 1): summary of results	121
Table 5.10 – Na ₂ SO ₄ crystallization tests (set 1): summary of results	121
Table 5.11 – Tests with pure water (set 1): summary of results	122
Table 5.12 – Relative equilibrium humidity of aqueous NaCl or Na2SO4 solutions	126

Chapter 6 – Case studies

Table 6.1 – Salvas rear façade: sampled materials	174
Table 6.2 – Salvas rear façade: IC on samples 0-2cm	175
Table 6.3 – Salvas rear façade: XDR on fine fraction of render	175
Table 6.4 – Tide mill NW wall: sampled materials	175
Table 6.5 – Tide mill NW wall: XDR on efflorescence	176
Table 6.6 – Despacho internal wall: Sampled materials	177
Table 6.7 – Despacho internal wall: XDR on efflorescence	178
Table 6.8 – Sta. Clara NW wall: Sampled materials	179
Table 6.9 – Sta. Clara NE wall: Sampled materials	180
Table 6.10 – Sta. Clara: XDR on efflorescence or sanded surface material	181
Table 6.11 – Sta. Clara: IC on samples 0-2cm	181
Table 6.12 – S. Sebastião exterior: sampled materials	182
Table 6.13 – S. Sebastião exterior: IC on samples 0-2cm	182
Table 6.14 – Rendering systems in the test panels	184
Table 6.15 – Summary of decay and diagnosis	189

Annex I – Magnetic resonance imaging

Table I.1 – Water content of MEP plaster and brick after total immersion in pure water	AI.5
Table I.2 – Water content of plaster L and stones D and M at capillary saturation	AI.8

Annex II – Making of the test specimens for set 1 tests

Table II.1 - 7	Tested specimens	All	.2
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Glossary of initials, acronyms and abbreviations

Initials and acronyms

ASTM	- ASTM International, originally known as the American Society for Testing and Materials
BRE	- Building Research Establishment
CEN	- European Committee for Standardization
COMPASS	- Compatibility of plasters and renders with salt-loaded substrates in historic buildings (European research project, EC contract EVK4-CT-2001-00047)
EC	- European Commission
EN	- European standard
FCT	- Fundação para a Ciência e a Tecnologia (Foundation for Science and Technology)
IET	 Instituto de Ciencias de la Construcción Eduardo Torroja (Eduardo Torroja Institute of Construction Sciences)
IPPAR	 Instituto Português do Património Arquitectónico (Portuguese Institute of Architectural Heritage)
ISO	- International Organization for Standardization
LNEC	- Laboratório Nacional de Engenharia Civil (National Laboratory for Civil Engineering)
LRMH	- Laboratoire de Recherche des Monuments Historiques (Historical Monuments Research Laboratory)
RILEM	- International Union of Laboratories and Experts in Construction Materials, Systems and Structures
TNO	- The Netherlands Organisation for Applied Scientific Research
TUE	- Technical University of Eindhoven
UNI	- Italian Organization for Standardization
WTA	 Wissenschaftlich-Technischen Arbeitsgemeinschaft f ür Bauwerks- erhaltung und Denkmalpflege (International Association for Science and Technology of Building Maintenance and Monuments Preservation)

Abbreviations

%V	- percent by volume
%W	- percent by mass
1D	- one-dimensional
2D	- two-dimensional
a.u.	- arbitrary units
CS	- cross-section
FID	- free induction decay
HMC	- hygroscopic moisture content
IC	- ion chromatography
m	- molal (mol/kg)
MC	- moisture content
MIP	- mercury intrusion porosimetry
MRI	- magnetic resonance imaging
NMR	- nuclear magnetic resonance
RF	- radio frequency
RH	- relative humidity
RH_{eq}	- relative equilibrium humidity of a salt solution
RH _{eq} ^{sat}	- relative equilibrium humidity of a salt; corresponds to the relative equilibrium humidity of a saturated solution of that salt
S/N	- signal/noise ratio in MRI techniques
Stage I	- first stage of the drying process of a porous material
Stage II	- second stage of the drying process of a porous material
Stage III	- third stage of the drying process of a porous material
Т	- temperature
WAC	- water absorption coefficient
XRD	- X-ray diffraction

Glossary of Symbols

<u>Latin</u>

а	- amount of hygroscopic moisture absorbed, in equilibrium conditions, by a sample of salt with unitary mass
A	- water absorption coefficient (WAC)
a_w	- water activity
b	- amount of hygroscopic moisture absorbed, in equilibrium conditions, by a sample of a porous material with unitary mass
В	- capillary penetration coefficient
B ₀	- strength of the main magnetic field used in NMR techniques
B ₁	- secondary oscillating magnetic used in NMR techniques
с	- solute concentration in a supersaturated solution
Cs	- solute concentration in a saturated solution
C_{v}	- concentration of water vapour
d	- penetration of the meniscus in a capillary
D	- coefficient of ion diffusivity in a porous medium
dr	- drying rate
D_v	- diffusion coefficient or diffusivity (of water vapour in a porous material)
E	- HMC of a salt at the set-point RH in a climatic chamber
E	- HMC of a salt at the minimum RH attained in a climatic chamber
E ⁺	- HMC of a salt at the maximum RH attained in a climatic chamber
f	- Larmor frequency; resonance frequency of nuclei in NMR techniques
G	- magnetic gradient used in NMR techniques

g	- gravitational constant
G _x	- x-axis component of G
G_y	- y-axis component of G
h	- hydraulic head
Н	- height of capillary rise
H _o	- height of capillary rise in equilibrium conditions in a cylindrical capillary
НМС	- HMC of a sample; given as a percentage by weight
HMC⁻	- HMC of a sample at the minimum RH attained in a climatic chamber
HMC^{+}	- HMC of a sample at the maximum RH attained in a climatic chamber
HMC ^e	- HMC of a sample at the set-point RH in a climatic chamber
j	- water vapour diffusion flux across a porous material
K ₁	- unsaturated permeability coefficient or liquid conductivity (across a porous material)
L	- length
1	- distance
т	 molality of the solution (concentration expressed in number of moles of solute per litre of solvent)
М	- molar mass
\overrightarrow{M}	- net magnetization; vector sum of the magnetic moments of all protons in a sample
m _{dry}	- dry mass
mi	- mass of a sample at a given weighing <i>i</i>
<i>m_{mat}</i>	- dry mass of a sample of a porous building material
m _s	- mass
<i>m_{salt}</i>	- dry mass of a salt sample
<i>M_{salt}</i>	- molar mass of a salt
M_w	- molar mass of pure water
p	- water column weight
p _c	- capillary suction, capillary pressure or suction stress
p _{cr}	- pressure on the loaded face of a growing crystal

Pe	- Péclet number
p_l	- ambient pressure
p_{v}	- partial pressure of water vapour in air
p _{vi} , p _{ve}	 partial pressures of water vapour in air at the two sides (interior and exterior) of a given element
Q	- amount of hygroscopic moisture in a sample of salt or porous material
Q_{eq}	- amount of hygroscopic moisture that forms a saturated salt solution
r	- radius of capillary
R	 ideal gas constant (8.3145 MPa.cm³.mol⁻¹.K⁻¹)
RH	- minimum RH in a climatic chamber
RH^{+}	- maximum RH in a climatic chamber
RH ^e	- set-point RH in a climatic chamber
S	- amplitude of the NMR spin-echo signal
s'	- thickness of air layer
t	- time
Т	- absolute temperature
<i>T</i> ₁	- spin-lattice relaxation time in NMR techniques
<i>T</i> ₂	- spin-spin relaxation time in NMR techniques
TE	- echo-time in NMR techniques
TR	- repetition time in NMR techniques
u	- density of the liquid flux across a porous material
Vc	- molar volume of a solid crystal
W	- total amount of liquid water absorbed by a porous material
W ₀	- positive intercept of the absorption curve of a porous material on the Y axis
x	- salt content in a sample; given as a percentage by weight
X _{crit}	- maximum salt content in a sample for which the mass variation of the sample due to fluctuation of the RH in a climatic chamber is equal to a percentage β of the dry mass of the sample

z - gravitic potential or distance to the ground

<u>Greek</u>

α	 parameter expressing the relevance of the possible error due to the uncertainty associated to HMC measurements in a climatic chamber or salt content estimates by the HMC method
α (Annex I)	- radial axis of changeable direction; radial profiles are measured in the direction of α by the 2D MRI technique
β	- percentage of the dry mass of a sample used as criterion for considering that equilibrium was achieved, typically, within a drying process
γ	- gyromagnetic ratio of nuclei
δ	- diffusion coefficient of water vapour in air
Δm	- mass variation between consecutive weighings of a sample
Δρ	- crystallization pressure
Δx	- uncertainty associated to the measurement of the salt content <i>x</i> in a sample
η	- dynamic viscosity of water
θ	- water content
θ_m	- water content at capillary saturation
μ	- water vapour resistance of a porous material
ν	- stoichiometric parameter; corresponds to the number of moles of ions produced when a mole of solute molecules is dissociated
П	- water vapour permeability of a porous material
П ^{аir}	- diffusion coefficient of water vapour in air
ρ	- bulk density of water
ho (Annex I)	- density of the nuclei in a sample subjected to NMR measurements
σ	- surface tension of a liquid
φ	- contact angle liquid / pore wall

 φ (Annex I) - rotation angle for measuring radial profiles by the 2D MRI technique

- ϕ osmotic coefficient
- ψ capillary potential

Chapter 1 – Introduction

"A experiência é a madre das coisas e por ela soubemos radicalmente a verdade" (Because experience is the mother of things, by it we discovered the very root of truth) Duarte Pacheco Pereira, in *Esmeraldo de situ orbis*, 1505-1508.

1.1 Motivation and scope

Soluble salts can cause major damage to porous materials in man-made constructions and geological outcrops. In buildings, soluble salts cause aesthetical problems, degrade health conditions in rooms and often require high-cost recurrent repairs. In extreme situations, soluble salts may endanger the structural safety of old constructions. A progressive and widespread loss of historic material is another, unfortunately common, consequence.



Fig. 1.1 - Two typical salt-damage patterns. Left: efflorescence and damage to paint, temporary exhibition gallery, south wing of Alcobaça Monastery, 2002. Right: sanding of lime plaster and of the lime bedding mortar leading to masonry disintegration, Paulistas tide-mill, Corroios, 2004.

Salt damage (Fig. 1.1) originates from ions (chloride, sulfate, nitrate or others) that migrate, dissolved in liquid water, in the pore network of building materials. Salt crystallization can occur mostly as a result of evaporative processes or temperature changes that cause these solutions to supersaturate.

Salt decay is today considered one of the main weathering processes of porous construction materials (Goudie and Viles 1997, Harris 2001, Scherer 2006a) and it is especially relevant in old buildings, i.e., those built before the use of Portland cement. Several factors can account for this particular relevance of salt decay in old buildings (Gonçalves 2003b, for instance):

- <u>Salts and water</u> Soluble salts are commonly more active in old buildings because:

 most of these buildings are based on thick solid walls made of porous hydrophilic materials that favour the ingress and permanence of water and salt solutions in their pore network; (ii) the walls were normally built in direct contact with the ground, thus, rising damp is a common feature; (iii) salts and moisture from different origins often accumulate in the masonry over the years, namely during periods of constructive deterioration (roof damage, masonry cracking, etc.).
- <u>Physical medium</u> Salt decay progresses faster in old masonry because these are currently based on lime mortars which are mechanically weaker than the porous materials in new buildings.
- <u>Structural impact</u> Salt damage may have more serious consequences in old buildings where the walls often have a structural function and, hence, masonry damage may eventually lead to structural collapse of the building.
- <u>Historic relevance</u> Many old buildings have a high historic value. Hence, whether they
 are single monuments or part of architectural assemblies, they should be preserved for
 future generations, as stated in the "Venice Charter" (ICATHM 1964). In contrast with
 current buildings, the concept of service life does not therefore apply to historic
 buildings. They should be preserved into a non-dated future.
- <u>Artistic relevance</u> Old buildings often include valuable sculptures or other artistic elements that can be lost or damaged by soluble salts.

Evaporation-induced crystallization is probably the most common case within salt decay processes. This is why plasters and renders play a key role in these processes. In fact:

- Plasters and renders are the surface layers of walls. They condition the exchange of
 moisture between the masonry and the environment and, hence, strongly influence
 moisture migration and drying processes in the entire masonry. Therefore, they may
 determine salt decay processes and features, as will be discussed in this thesis.
- Along with a sanitary or aesthetic role, plasters and renders had (and have) a main sacrificial role: they protect the masonry from destructive actions such as those from wind, rain, human action or salt crystallization. Indeed, plaster/render damage is not as harmful as (structural) masonry damage. Furthermore, it is much easier to repair. Because damage by salt crystallization usually starts at the surface of walls and progresses as the superficial layers are destroyed, plasters and renders minimize the occurrence of masonry damage. This is probably one of the reasons why, in many countries such as Portugal, plasters and renders were normally used on (weaker) ordinary masonry, currently made from irregular stone elements and lime-based bedding mortar (Leitão 1896), and less used on (stronger) ashlar masonry. Segurado (1908), for instance, argues that plastering/rendering of ordinary masonry is indispensable.

Due to the high number of buildings affected by salt decay features, as well as to the harmfulness and complexity of such processes, more and more researches are carried out in the subject involving chemists, physicists, geologists and engineers. However, despite of many advancements made, several fundamental aspects of salt decay processes are not yet fully understood, more specifically, the (micro-scale) damage mechanisms, salt-induced dampness or the (macro-scale) behaviour of building elements. Accordingly, adequate strategies of prevention or mitigation are often lacking (Harris 2001, Scherer 2006a).
The behaviour of plasters and renders in relation to salt crystallization is one of the topics where further research was needed. In Portugal, for example, end-users complain that salt decay is often aggravated after restoration interventions on plasters/renders. Further, despite their relevance in salt decay processes, no standard crystallization tests exist for plasters and renders, in contrast to other materials such as stone or brick. The need for research on this topic justified the launch of the European project, COMPASS - Compatibility of plasters and renders with salt-loaded substrates in historic buildings. Some of the work presented in this thesis, namely in sections 3.4, 3.5, 5.5 and 6.2 to 6.4, was carried by the author under the framework of the COMPASS project.

1.2 Aims and objectives

This thesis aims at understanding the behaviour of plasters and renders on salt-loaded walls. Its main objectives are the following:

- To assess the possibilities and limitations of two general laboratory techniques, for relative humidity (RH) control using salt solutions and for salt content determination by means of hygroscopic moisture content (HMC) measurements, in order to decide whether and in which conditions they can be adequate experimental tools.
- To gain further understanding of the drying features of salt-loaded masonry, namely, the influence that soluble salts have on drying of porous materials, on salt distribution during drying and on salt-induced damp mechanisms.
- To investigate the possible influence that factors, such as the type of salt, presence of a paint layer, type of substrate material, use of water-repellent additives or the material saturation conditions have on the behaviour of plasters and renders in relation to salt crystallization.
- To assess the possibilities and limitations of a given crystallization test as a performance test for plasters and renders.
- To identify factors that may account for sodium chloride, which is frequently involved in real decay processes but in laboratory crystallization tests is normally much less damaging, or even completely innocuous, than salts such as sodium sulfate.
- To understand the nature and causes of some serious salt decay problems in old Portuguese buildings, specifically as regards the type of salts and sources of moisture.
- To compare the possibilities and limitations of expertise-based diagnostic methodologies, that rely mostly on visual inspections, and those based on physical/chemical testing.
- To test the application technique and performance of three industrial plasters available on the Portuguese market for salt-loaded walls.
- To identify chief factors that can account for the worsening of dampness and decay problems after restoration interventions on plastered/rendered buildings
- To provide guidelines for selecting plasters and renders for moist salt-loaded walls.

1.3 Outline of the thesis

This thesis was structured into seven different chapters and three annexes. The first chapter is the present introductory chapter. The remaining six chapters and annexes address the following subjects:

- Chapter 2 concerns the current state-of-the-art and state-of-the-practice in salt decay pathology and mitigation processes. There is particular focus on old plastered/rendered buildings and on the situation in practice in Portugal. The state-of-the-art is assessed by means of a review of published scientific literature. The state-of-the-practice is approached mostly on the basis of interviews with Portuguese end-users and of a survey of the Portuguese market regarding industrial products for moist salt-loaded walls.
- In Chapter 3 the possibilities and limitations of two general laboratory techniques are evaluated. The method of RH control by salt solutions is discussed on the basis of experimental research on sodium chloride. The HMC method for salt content determination is assessed by means of both theoretical and experimental research on nine different salts.
- Chapter 4 concerns the drying behaviour of salt-loaded materials. This research is mostly based on drying experiments monitored by a 2D (two-dimensional) MRI (magnetic resonance imaging) technique.
- Chapter 5 discusses the behaviour of plasters and renders in relation to salt crystallization. Five different sets of crystallization tests or drying tests monitored by the same 2D or by a 1D (one-dimensional) MRI technique are presented and serve as a basis for: (i) assessing the influence of various factors such as the type of salt, nature of the substrate material, presence of a paint layer, material saturation condition or use of mass water-repellent additives (ii) identifying different factors that can account for an aggravation of salt decay problems after restoration interventions; (iii) examining the possibilities and limitations of using salt crystallization tests to predict the performance of plasters and renders on saltloaded walls; (iv) discussing how to choose plasters and renders for salt-loaded walls.
- Chapter 6 presents the five case studies investigated within this thesis: Salvas Chapel (Sines), Alhos-Vedros tide-mill (Moita), Despacho House (Pereira), Cloister of Sta. Clara-a-Nova Monastery (Coimbra) and São Sebastião Church (Almada).

Site inspections and a systematic sampling/testing procedure were used to assess the nature and causes of the decay in these buildings. On the basis of this field research, some factors, additional to those discussed in Chapter 5, that may account for an aggravation of salt damage after restoration interventions are discussed. Further, a comparison between expertise-based and sampling/testing-based diagnostic methodologies is made. Finally, the application technique and performance of industrial rendering systems for salt-loaded walls are evaluated by means of test panels made in São Sebastião church.

• Chapter 7 presents the overall conclusions achieved, focusing on the aims and objectives of the thesis. A summary of the work performed and the main topics on which further research is needed are also presented in this final chapter.

- In chapters 3, 4 and 5 background information on specific topics such as hygroscopicity, moisture transport, sorption and drying of porous materials or salt crystallization tests is presented to allow a better understanding of the main subjects addressed in each chapter.
- Annex I introduces the principles of MRI and presents the 1D and 2D techniques used.
- Annex II describes the making of the test specimens used in set 1 tests of Chapter 5.
- Annex III includes the damage assessment and diagnosis form used for guiding the inspections of the case studies presented in Chapter 6.
- A glossary of initials, acronyms and abbreviations and a glossary of symbols are included in the thesis immediately before the present chapter.

1.4 Delimitations of scope and key-assumptions

This thesis deals with salt decay in plasters and renders. Plasters and renders are here defined as mortar-based coverings used on the interior and exterior, respectively, of buildings. Even in the cases where, as a simplification, only plasters are mentioned, all the work presented in the thesis concerns and is applicable to both types of coverings.

Salt decay is a weathering process arising from the presence of salt crystals on the surface of porous materials (efflorescence) or from mechanical stresses introduced by salt crystals deposited in the material pores (subflorescence). In a broad sense, salt decay may also refer to salt-induced dampness features. Other processes, such as those primarily based on chemical reactions between soluble salts and building materials, are not covered. In this thesis, "salt damage", "damage by soluble salts", "damage by salt crystallization", "salt weathering" or related terms refer to salt decay processes, as defined above.

Damage by soluble salts is particularly current and harmful in old buildings, as stated in section 1.1. Therefore, although many topics and conclusions in this thesis are also applicable to recent buildings, by default, the case of old buildings is addressed.

Salt decay requires the simultaneous presence of soluble salts and moisture in the porous material. Although this moisture may have different origins (Massari and Massari 1993, for instance), rising damp is probably the most frequent and perhaps one of the hardest to solve sources of moisture in old buildings. Hence, in this thesis, rising damp is considered the default case. Unless otherwise specified, moisture is assumed to be due to rising damp.

Salt crystallization may arise from temperature changes for salts whose solubility varies with temperature. It may also arise from the contact between the salt solution and pre-existing salt crystals when these correspond to phases with different solubility. As will be explained in the following chapters, both processes may take place, for instance, with sodium sulfate. They can be very relevant in laboratory tests where the material is subjected to high temperatures or wet-dry cycles ranging from moistness to dryness. In buildings, however, evaporative crystallization is likely to be the most relevant process. Therefore, in this thesis, this will be the mechanism considered by default and focused in greater detail.

Salt ions generaly migrate in porous materials by advection, that is, carried by the flowing liquid water. Yet, ion diffusion in the liquid phase can also take place due to gradients of ion concentration. For instance in evaporative processes, ion concentration increases at the drying front and, hence, a backwards migration of salt can arise which tends to minimize the concentration gradient. As it will be explained latter, ion diffusion can be relevant for very low liquid fluxes. In the experiments in this thesis, ion diffusion will not in general be considered due to the short time periods involved and because no evidence was found that it had any relevancy.

Chapter 2 – Research and practice: current situation

2.1 Introduction

Since the first crystallization tests on stone carried out by Brard for simulating the disruptive action of ice (Thury 1828) salt crystallization has been progressively recognized, at a scientific level, as a specific damage process. Today it is considered one of the most powerful decay mechanisms of porous materials in man-made constructions and geological outcrops.

Salt decay has been attracting more and more attention from the scientific community, especially over recent decades, as demonstrated by the growing number of research articles and by some general reviews on the subject (Evans 1969-1970, Charola 2000, Doehne 2003, Rodrigues and Gonçalves 2005). But, despite the great deal of progress made, it is still one of the least understood deterioration mechanisms of porous building materials (Rodrigues 1991, Scherer 2006a).

Despite the lack of comprehensive scientific explanations, salt decay is something that practitioners, particularly those working with old buildings, have to deal with currently. In fact, damage by salt crystallization has long been known, as shown by Herodotus' (484 B.C. - 425 B.C.) observation that "salt exuded from the soil to such an extent as even to injure the pyramids". In many constructions, especially those composed of weaker masonry of lime mortar and stone or brick, plasters and renders were traditionally used as sacrificial layers for protection against damaging actions, namely, salt crystallization. Salt decay was typically handled by means of assiduous maintenance that included repainting, localized repairs and periodic replacement of plasters and renders.

This situation progressively changed after the industrial revolution. New materials appeared which behave differently when used in moist and salt-loaded walls, namely cement-based mortars and emulsion paints. More demanding, health and aesthetical, requirements arose both from higher living standards and from the increasing interest in the conservation of old buildings. Decay caused by salts from cement-based materials, conservation treatments or air pollution became often more relevant than that caused by salts from natural sources such as the soil or seawater. In fact, although the care of old buildings was one of the driving motivations for scientific research in the field (Jokilehto 1999), conservation interventions were eventually found to be a frequent cause of damage. Further, severity of material durability requirements increased not only because better performances were expected by

more demanding end-users but also because of the higher labour costs associated with repair and maintenance work. Traditional knowledge in construction techniques could no longer respond to the new situations and was gradually lost from generation to generation.

Today, as scientific knowledge on salt decay slowly progresses, end-users are forced to solve practical decay problems. The building material industry, through a combined scientific and empirical approach, tries to respond to the needs of end-users. Special plasters and renders have been developed that are aimed at being more durable or at presenting specific performance features, such as preventing the occurrence of surface damage.

This chapter is aimed at giving an overview of the present situation of salt decay theory and practice focusing particularly on the case of plasters and renders for old buildings. The state-of-the-art is discussed in section 2.2 and the state-of-the-practice in section 2.3.

2.2 State of the art

2.2.1 - Salt decay pathology

Salt decay requires the simultaneous presence of soluble salts and water in the pores of building materials, as well as appropriate environmental conditions. Although these deterioration processes are very complex, the general pathology can be described (Fig. 2.1):



Fig. 2.1 - Salt decay pathology

There are three main types of factors underlying salt decay processes as seen in Fig 2.1: (i) the predisposing factors, soluble salts and moisture, without which damage cannot occur; (ii) a physical medium, that is, the porous material where the process takes place; (iii) adjuvant factors which condition the occurrence and influence the process, thereby determining the damage patterns.

The cause of salt decay, which arises from the joint influence of these three types of factors, is the occurrence of salt crystals either on top (efflorescence) or inside (subflorescence) the porous material. In evaporative crystallization processes, crystals precipitate at the drying front where the solute evaporates and, hence, the solution concentration increases up to (or above) the maximum solubility of the salt. In such processes, the drying front tends to find a position that corresponds to equality between the liquid and vapour fluxes that cross the material to and from that front, respectively. Efflorescence occurs when the liquid flux is high enough to compensate the evaporative demand. In that case, the liquid is able to reach and evaporate on the outer surface of the material. Subflorescence takes place when the liquid flux is lower than this and, hence, the drying front is located inside the material.

Building anomalies caused by salt crystallization can manifest as a wide range of signs (Fig. 2.1). These anomalies can configure three different types of damage: material damage, aesthetical damage or damage associated to the presence of moisture.

Subflorescence can induce internal stresses that overcome the mechanical strength of the porous material and, hence, cause it physical damage. Efflorescence does not constitute or cause material damage, though it may account for aesthetical or dampness problems. However, efflorescence can be redissolved, then reabsorbed into the material and eventually recrystallize as subflorescence.

2.2.2 - Signs

Common damage patterns of porous building materials are described in Normal 1/88 (ICR and CNR 1988) which was recently re-published as UNI Standard 11182 (UNI 2006). A "damage atlas" focusing specifically on damage patterns of plasters and renders was drawn up within the COMPASS project (TNO 2002).

Some examples of common salt decay patterns are shown below in Fig 2.2. These were observed on traditional plastering / rendering systems during preliminary inspections carried out within this thesis on 31 buildings and monuments. Other patterns will be presented in Chapter 6 when describing the five case-studies selected out of these 31.

The damage patterns in Fig. 2.2 occur when soluble salts crystallize on or close to the surface of the wall and, thereby producing efflorescence, dampness problems or physical damage to the surface layer of the plaster or render. But salts may also deposit deeper in the wall. In this case, damage may arise without being immediately visible as, for instance, detachment of plaster/render coats from each other or from the substrate. This sort of damage can often be detected, even while there are no visible signs of it, simply by tapping the surface: a hollow sound indicates that detachment has occurred.



(a) Efflorescence – New See basement, Coimbra, 2003



(b) Salt-damp – Cloister of Sta. Cruz Monastery, Coimbra, 2002



(c) Salt damp and staining – Alcobaça Monastery, 2002



(d) Paint peeling – building in the historic centre of Santarém, 2003



 (e) Scaling of lime-wash / lime-render system – Torre das Cabaças, Santarém, 2002



 (f) Delamination (layering of material with laminated structure, in this case, a render applied in different coats)
 Tibães Monastery, Braga, 2002



(g) Exfoliation (layering of material with non-laminated structure, in this case, the surface coat of the render) – Sta. Clara-a-Nova Monastery, Coimbra, 2005



(h) Sanding of render – Salvas Church, Sines, 2003



(i) Erosion – Arouca city hall library, 2002

Fig. 2.2 – Some common salt decay patterns of plasters and renders. Damage pattern definition according to the Plaster Damage Atlas (TNO 2005). When soluble salts crystallize in the pores of building materials they can have a cementing rather than a disruptive effect (Rossi-Manaresi and Tucci 1991). In this case, damage is not likely to arise straight away. However, the hardened surface layer tends to behave differentially, namely under thermal or hydric dilation processes, and can eventually detach.

Alveolar (or honeycomb) decay is a particular pattern where numerous small cavities separated by hardened walls develop on the surface of the material. Though this sort of decay is also observed in hot deserts or Arctic landscapes, it seems to develop especially well in costal areas, particularly on the Atlantic coast (Goudie and Viles 1997, for instance) and on the spray zone above ordinary wave action level (Rodríguez-Navarro *et al.* 1999). Alveolar decay is typically observed in stone materials but it can also occur in rendering mortars, as seen in Fig. 2.3. However, despite its scientific interest, this type of decay will not be focused on this thesis because it is associated with long time periods. In fact, in contrast to stone materials, one of the main features of plasters and renders is their replaceability. Hence, though alveolar decay may be significant in very specific cases (for instance, archaeological ruins), it is not expected to be relevant in buildings subjected to periodic maintenance.

In plastered/rendered walls, salt decay patterns are typically observed on the plaster or render but, especially when those have already suffered extensive damage, can affect the masonry as well, as seen in the examples of Figs. 2.3 and 2.4.



Fig. 2.3 - Alveolar decay – building adjacent to a branch of the Tagus river, at less than 10 km from the open sea, Alhos-Vedros, 2002



Fig. 2.4 - Masonry disintegration - Paulistas tide mill, Corroios, 2004

2.2.3 - Moisture

Soluble salts originate from ions that are transported while dissolved in liquid water that migrates in the pore network of building materials. Liquid water can penetrate into these materials by different processes (Paiva 1969, Massari and Massari 1993, Henriques 1994, for instance):

 Construction humidity originates from the water used in the mixing of mortars or setting of bricks, from the use of materials previously exposed to rain or from specific processes such as water-jet cleaning.

- Rising damp is ground moisture that rises up in the walls by capillarity. It can be either phreatic water or rain water that penetrated into the soil surface layers.
- Dew point condensation is another possible source of moisture. When humid air comes into contact with colder surfaces its temperature drops and, in consequence, its RH increases. If the absolute saturation humidity is exceeded, dew point condensation can occur on the cold surface. Dew point condensation may also take place inside masonry materials. Since the absolute saturation humidity varies with temperature, it may be attained somewhere across a material subjected to a temperature gradient.
- Hygroscopic moisture is attracted from the air. Both common building materials (such as mortars, brick or stone) and soluble salts have hygroscopic properties. However, soluble salts are much more hygroscopic. Accordingly, hygroscopic dampness can be particularly severe when salts are present. The mechanisms of hygroscopicity will be discussed in more detail throughout this thesis.
- Penetration of rainwater can take place through cracks, construction joints, damaged roofs, etc.
- Moisture can also arise from accidental causes such as leaking of pipes or fouling of gutters.

Liquid water is ordinarily present in old building materials. In fact, while in new buildings dampness is typically acute but transitory and results mostly from poor construction, in old buildings dampness is chronic by nature (Massari and Massari 1993). The major source of moisture in these buildings is probably rising damp because, as pointed in the previous chapter, old buildings are ordinarily based on thick solid walls built with porous hydrophilic materials in direct contact with the ground. Further, old buildings often went through periods of constructive deterioration that may have allowed penetration and further accumulation in the masonry of moisture and salts. And, since soluble salts are common in these buildings, hygroscopic damp occurs often. As for dew point condensation, its relevance as respects salt decay processes in old buildings is mostly unknown.

2.2.4 -Soluble salts

Old masonry normally contains soluble salts. A large variety of salts can be found in these walls, namely chlorides, sulfates, nitrates and carbonates of sodium, potassium, magnesium or calcium (Arnold 1981). According to Goudie and Viles (1997), sodium chloride, sodium sulfate, sodium carbonates and gypsum are particularly common.

Soluble salts can have different origins. In some cases they may have been involved in complex chemical reactions (Arnold 1981) that impede tracing their sources. Nonetheless, the type of salt is still the best indicator of its origin. Sodium chloride, for instance, often arises from direct contact of the building materials with seawater, by marine fog, seawater spray or contaminated groundwater. Sodium chloride may also come from salty goods once stored in the building or even, due to human consumption of sodium chloride, from

groundwater contaminated by domestic residues (Arnold and Zehnder 1989). Groundwater often carries nitrates produced by decomposition of organic materials, particularly in soils treated with organic fertilizers. Nitrates may also be introduced by animal excrements and tissues, for instance in buildings used as stables or colonized by pigeons, or by microbiological activity in building materials. Sulfates, for instance of sodium, can also originate from the soil (Schaffer 1932). Carbonate salts typically originate from high alkali content materials, for example, cement-based mortars (Arnold 1981). The alkali hydroxides in these materials react with carbonic acid (which results from dissolution of air CO₂ in the water in damp walls) and thereby transform into alkali carbonates. In polluted environments, the alkali hydroxides (Arnold 1981) or the resulting alkali carbonates (Schaffer 1932) may eventually form alkali sulfates by reaction with sulphuric acid from the atmosphere. Ceramic materials may also carry salts into the masonry, namely sulfates of sodium or magnesium (Schaffer 1932). The gypsum that composes many plasters may dissolve and migrate to other masonry materials. Gypsum may as well form, in polluted atmospheres, by reaction of sulphuric anhydride (SO₃) in the air with calcium carbonate in calcareous stones or lime mortars. Further, many other salts and salt origins are possible in old buildings.

Soluble salts crystallize mostly as a consequence of evaporative processes or temperature changes. In this context, the two fundamental characteristics of soluble salts are their solubility and relative equilibrium humidity.

Solubility is the ability of a salt to dissolve, in this case, in water. It may be expressed quantitatively as the maximum amount of salt that can be dissolved at a certain temperature. When the water contained in a dilute salt solution evaporates, the concentration of the solution increases. If the concentration that corresponds to the maximum solubility (or simply "solubility") of the salt is exceeded, the salt in excess can crystallize. Crystallization may also arise from temperature changes in the case of salts whose solubility varies with temperature. In both cases, crystallization may not occur readily because salt solutions can supersaturate, that is, attain non-equilibrium states where a higher amount of solute is dissolved than that which can be held in equilibrium conditions. But all systems tend to equilibrium and, hence, crystallization will ultimately take place. The higher the supersaturation of the solution the higher the tendency of the salt to crystallize. The maximum supersaturation that salt solutions can reach under certain conditions is called the metastable limit (Strege 2004, for instance). At this point crystallization occurs readily.

Salt crystallization from solution proceeds in two main phases. In the first phase, the nucleation phase, very small crystals are formed, often randomly. These nuclei grow during the subsequent crystal growth phase. Nucleation occurs more easily in the presence of the so-called "seeds" (for instance, solid dust particles) than in homogenous solutions. Therefore, the presence of seeds may strongly condition the ability of a salt solution to supersaturate.

The relative equilibrium humidity (RH_{eq}^{sat}) of a salt corresponds to the RH below which the solid crystal is in equilibrium with air at a certain temperature. When the environmental RH attains this value, the crystal attracts moisture from the air and ultimately dissolves forming a saturated solution. Above the RH_{eq}^{sat} of the salt, the higher the air RH the more dilute the resulting salt solution. At 100% RH, salt solutions tend to become infinitely diluted.

Conversely, when a salt solution is exposed to an RH lower than the RH_{eq}^{sat} of the salt, all the salt in that solution eventually crystallizes while the solvent (water) is evaporated.

Many soluble salts possess different equilibrium crystal phases for distinct conditions of temperature or RH. Some salts can even exist in non-equilibrium (metastable) crystalline phases which, although they tend to transform into the equilibrium phase at that temperature and RH, can be stable for long periods.

The different phases of a salt or salt solution and the conditions under which each phase exists are expressed in the so-called phase diagrams. In such diagrams, lines of equilibrium (phase boundaries) are used to represent the phase transitions that the substance may undergo. Phase diagrams can address numerous types of parameters and properties. In the present case, there are two types of phase diagrams that can be of interest: (i) solubility phase diagrams (Figs. 2.6 and 2.8) where the solution concentration is plotted against the temperature; phase boundaries are used to represent the solubility of the possible crystal phases; (ii) RH-temperature phase diagrams (Figs. 2.7 and 2.9) that express the environmental conditions under which the salt is stable as a certain crystalline phase or as solution; phase boundaries represent the transition between the possible phases. Strictly, phase diagrams concern equilibrium situations. Yet, for practical reasons, non-equilibrium features are often included, usually as dashed lines, such as the solubility of metastable forms or supersaturation boundaries.

Salt crystals may present different shapes, that is, different habits. They can be acicular (needle-like crystals), fibrous (hair-like crystals), equant (with nearly equal dimensions), etc. Habits do not depend exclusively on the internal structure of the crystal. They are influenced also by the conditions under which the crystal grows, for instance the environmental conditions (Bloss 1971, for example).

When a salt is dissolved in water it forms an electrolyte solution or, simply, electrolyte. An electrolyte contains free ions and is able to conduct an electric current. Indeed, when a soluble salt is mixed in water, its ions are chemically dissociated because water molecules have stronger (covalent) bonds than the salt molecules (ionic) bonds.

Building materials rarely contain a single type of salt. Normally a mixture of different salts occurs. The properties of mixed salt solutions cannot generally be predicted by weighing the properties of the individual salts due to the mutual interaction between the dissolved ions. Only in very dilute solutions can that interaction be neglected. The RH_{eq}^{sat} of salt mixtures, for instance, arises not as a single value but as a range of values that do not necessarily fall within the limits of the RH_{eq}^{sat} of the individual salts (Price and Brimblecombe 1994).

Also the solubility of the salts may change in the presence of other salts. As summarized by Charola (2000), in general: (i) if two salts do not have any ions in common, the solubility of both salts increases due to the higher ionic strength of the solution; the less soluble salt has a higher solubility increase; (ii) if two salts have an ion in common, their solubility decreases; the less soluble salt has a higher solubility decrease.

Sodium sulfate and sodium chloride are amongst the most common salts in decayed buildings. Moreover, these are the salts generally used in laboratory crystallization tests. Since they were also the salts used in the experiments for this thesis, a more detailed description of their properties will be presented here.

Sodium chloride (Figs 2.6 and 2.7) is perhaps the simplest and best known of all soluble salts involved in the decay of old buildings. Above 0°C, sodium chloride possesses only one crystalline form, the anhydrous halite (NaCl). The solubility of sodium chloride practically does not vary with the temperature. Normally, sodium chloride does not reach high supersaturations and, hence, it readily crystallizes at a concentration of around 6 molal.



Fig. 2.6 – Solubility diagram of sodium chloride. Data from Steiger (2006).



Fig. 2.7 – RH/T phase diagram of sodium chloride. Data from Steiger (2006).

Conversely, sodium sulfate (Figs. 2.8 and 2.9) is probably one of the most complex salts involved in salt decay processes. It can form three distinct crystalline phases, can easily supersaturate and its solubility is highly temperature dependent.

The stable phase in contact with a saturated solution of sodium sulfate is mirabilite $(Na_2SO_4.10H_2O)$ or thenardite (Na_2SO_4) for temperatures below or above 32.4°C, respectively. The heptahydrate $(Na_2SO_4.7H_2O)$ may also exist below 32.4°C in contact with a saturated solution but it is not an equilibrium form, that is, the solution is supersaturated with respect to mirabilite.

The RH-temperature diagram of sodium sulfate was recently (re)calculated by Linnow *et al.* (2006). As seen in Fig. 2.9, the only stable phase in the air above 32.4°C is thenardite. Below 32.4°C the three crystal phases are possible. For instance at the ordinary temperature of 20°C, thenardite is the stable phase below 77% RH (and not 71%, as previously accepted) and mirabilite is stable between 77% and 95.6% HR (and not between 71% and 93.6%, as previously accepted). Therefore, in drying processes, mirabilite can change into thenardite when the air RH is of 77% or less. Yet, thenardite solubilises only at 87% RH (and not 82%). Thus, in absorption processes, thenardite will probably only change into mirabilite when the air RH is of 87% or more. This is expected to happen because the hydration of sodium sulfate does not consist of direct absorption of moisture by the anhydrous phase but rather of dissolution of thenardite and its reprecipitation as mirabilite (Charola and Weber 1992, MacMahon *et al.* 1992, Rodriguez-Navarro *et al.* 2000). The heptahydrate can form between

77% and 95.6% RH but, since it is a metastable phase, it may transform into mirabilite at any time. The RH_{eq}^{sat} of mirabilite is 95.6% RH, therefore, in equilibrium conditions above this RH, a liquid solution of sodium sulfate exists.



2=solubility of the heptahydrate 3=solubility of thenardite 4=thenardite / mirabilite boundary

Fig. 2.8 – Solubility diagram of sodium sulfate. Adapted from Rodríguez-Navarro *et al.* (2000).



(a) deliquescence-crystallization of mirabilite
(b) hydration-dehydration
(c) deliquescence-crystallization of thenardite
Light grey - metastable branch



2.2.5 - Damage mechanisms

Soluble salts can cause three main types of damage to porous building materials: aesthetical damage, damage associated to the presence of moisture and material damage.

Aesthetical damage arises from the presence of efflorescence but can be associated with the two other types of damage, material damage and salt damp.

Dampness problems (Figs. 2.2 b and c) often arise from the presence of soluble salts in building materials. Although the mechanisms of salt damp are not fully understood (Harris 2001, for instance), hygroscopicity is generally pointed out as the cause of many of these problems (Massari and Massari 1993, Henriques 2006). In fact, the high hygroscopic absorption of soluble salts and the low RH_{eq}^{sat} of some salt species (for example, at 20°C, 75% RH for sodium chloride or even 33% RH for magnesium chloride) allow the absorption of high amounts of moisture from the air at relatively low RH.

Material damage, such as that in Figs. 2.2 d) to e), can occur when salts crystallize and thereby introduce internal stresses in the pore system of building materials. The following discussion in this and in the next section refers, by default, to this sort of damage.

It is generally accepted that material damage occurs when a confined crystal grows against the pore walls. This can happen when, due to repulsive forces between the crystal and the porous material, a film of solution interposes between the crystal and the pore wall. The higher the supersaturation ratio of this solution, the higher the pressure exerted by the growing crystal. This was first expressed by Correns' equation (Correns 1949):

$$\Delta p = p_{cr} - p_l = \frac{R T}{V_c} \ln\left(\frac{c}{c_s}\right)$$
 Eq. 2.1

In this equation, Δp (MPa) is the crystallization pressure, p_{cr} (MPa) the pressure on the loaded face of the growing crystal, p_i (MPa) the ambient pressure, T (K) the absolute temperature, V_c (cm³. mol⁻¹) the molar volume of the solid crystal, R (8.3145 MPa.cm³.mol⁻¹.K⁻¹) the ideal gas constant, c_s the solute concentration in a saturated solution and c the solute concentration in the supersaturated solution.

The subject of crystallization pressure is still largely under discussion. After Correns', other equations have been proposed, as summarized in the recent literature review by Steiger (2005a). The main objection to Correns' or similar theories comes from the fact that, in mechanical equilibrium conditions, when a film of solution surrounds the crystal, the pressure exerted on the pore walls depends on the curvature of the crystal. In that case, megapascal pressures sufficiently high to damage porous building materials could only develop in very small pores, smaller than around 100 nm (Scherer 2006a), which contradicts reality (Scherer 2004, Steiger 2005b). However, it is possible that Correns' theory might be applicable to non-equilibrium situations arising during drying processes where the liquid film is disrupted and some solution is trapped in the gap between the crystal and the pore wall, which then would act as a small pore (Scherer 2004).

Hydration pressure was for a long time considered a possible damage mechanism but some practical evidences contradict this possibility. Firstly, many salts such as sodium chloride do not possess hydrates in the normal temperature range. Secondly, hydration consists probably of dissolution of the anhydrous phase followed by reprecipitation of the hydrated phase, rather than of direct absorption of moisture by the anhydrous phase. This happens with sodium sulfate (Charola and Weber 1992, MacMahon *et al.* 1992, Rodriguez-Navarro *et al.* 2000), so, it is possible that the same occurs with other hydratable salts.

Other damage mechanisms were also considered to explain how salts damage porous building materials, as reviewed and discussed by Charola (2000). Hydrostatic crystallization pressure was once considered because the volume of a supersaturated solution is smaller than that of the resulting crystal plus saturated solution. But the solubility of salts increases with pressure. For this reason, it was argued that hydrostatic pressure could not generate pressures high enough to overcome the mechanical strength of most building materials. Some other damage mechanisms have also been proposed, such as shear stress by salt films on pore walls (Pühringer 1983) or ion diffusion (Lewin 1990), but their validity was not effectively demonstrated.

Soluble salts have a cementing rather than a disruptive effect when the crystallization pressure is lower than the mechanical strength of the material, as argued by Rossi-Manaresi and Tucci (1991). In that case, damage is likely to arise later due to differential (thermal or hydric) behaviour between the contaminated and the non-contaminated material.

The mechanism of hydric dilation, originally proposed by Snethlage and Wendler (1997), was the object of recent experimental research (Lubelli 2006). The author concluded that salt crystals (of NaCl, NaNO₃ or KCl) may form a layer on the pore walls and thereby induce relevant hydric dilation during drying. In contrast, non-contaminated materials expand during wetting. Hence, shear stresses due to differential hydric dilation could develop between the contaminated and the non-contaminated material and induce damage such as scaling, bulging or spalling. There is, however, some disagreement with this model. Scherer (2006b) argues that crystallization pressure rather than hydric dilation could cause the expansion of salt-loaded materials during drying. Some salt could crystallize on the walls of the larger pores as the (initially saturated) salt solution evaporated from these larger pores. If salt crystals were to grow in the smaller pores, high stresses would be generated which could totally disrupt the material. However, if a film of solution remains on the mineral surfaces, then the smaller crystals would be unstable with respect to the larger ones, so ions would diffuse through the film and attach themselves to the larger crystals. Since this process takes time, there would be a transient stress (until the small crystals had all dissolved) that could cause the observed expansion. If the film of solution becomes discontinuous, then the stress would continue to be exerted in the small pores. At the same time, a film isolated under the crust of salt on the walls of the large pores could contribute to the expansion.

The above considerations show that the mechanisms by which salts damage porous building materials are still far from established. Nonetheless, linear crystallization pressure seems to be today the most widely accepted mechanism (Scherer 2004, Steiger 2005a).

The occurrence of material damage and its patterns depend largely on the depth at which salts crystallize. In evaporative crystallization processes salts deposit within the pores when the liquid flux towards the drying surface is not high enough to compensate the evaporative demand, as explained in section 2.2.1. Lewin (1982) argued that the process of salt-decay "is characteristically manifested in the form of a thin layer (around 1 mm thick) of the surface that lifts up in the form of a blister, peels outward as a spall, flakes off, or powders away". Based on his own experimental work, he further postulated that "the necessary condition for the occurrence of this decay is the development of a steady state at the exposed surface, wherein the rate of evaporation of water via diffusion through a layer of the porous solid is balanced by the viscous flow of solution".

In this approach to salt decay mechanisms, ion transport is assumed to occur only by advection. Yet, in certain cases, ion diffusion may also play a role. The competition between advection and diffusion is expressed by the Péclet number *Pe.* For *Pe* << 1 diffusion dominates, whereas for *Pe* >> 1 advection dominates. In the case of a unidirectional drying experiment, *Pe* is given by Eq. 2.2:

$$Pe = \frac{dr.L}{\theta_m.D}$$
(Eq. 2.2)

In this equation, $dr (m^3.m^{-2}.s^{-1})$ is the drying rate, L (m) the length of the sample, $\theta_m (m^3/m^3)$ the maximum fluid content by capillary saturation and $D (m^2.s^{-1})$ the coefficient of ion diffusivity in the porous medium.

Equation 2.2 shows that the slower the transport of liquid towards the drying surface, the higher the significance of the (backwards) ion diffusion. This means that, at very low drying rates, ion diffusion may theoretically impede salt crystallization at the drying front.

2.2.6 - Influencing factors

Old masonry normally contains soluble salts and moisture. Although factors directly related to these predisponent factors are certainly relevant, such as the kind and quantity of salt and the amount of moisture, the occurrence and type of decay depends also, as depicted in Fig. 2.1, on other factors:

- Adjuvant factors, namely, the environmental conditions (air temperature, RH and velocity, solar radiation, etc.);
- Physical properties of the porous medium where the process takes place.

Salt decay processes are indeed complex and still largely unknown. Rodriguez-Navarro and Doehne (1999) argued that the intensity of damage and the damage patterns depend fundamentally on the solution supersaturation ratio and on the location of crystallization within the porous material. From these authors' and others' experimental work, several influencing factors could be identified that are listed and discussed bellow. Most of those works were carried on sodium chloride and/or sodium sulfate. Indeed, one of the issues currently addressed in salt decay researches is the behaviour of sodium chloride versus that of sodium sulfate. The reason for this interest is that, in laboratory crystallization tests, sodium sulfate typically causes extensive damage to porous materials while sodium chloride is often much less damaging or even completely innocuous. And, yet, sodium chloride is often found to be involved in salt decay processes in real constructions.

Solution composition

The velocity of capillary transport in porous materials may differ for distinct liquids due to changes in their viscosity, surface tension or contact angle with the material. Rodriguez--Navarro and Doehne (1999) sustain that, depending on the type of salt and solution concentration, the capillary flow can be relevantly slower for some salt solutions (for example, sodium sulfate in comparison to sodium chloride). Therefore, these solutions have more difficulty in reaching the drying surface and, hence, a higher tendency to form harmful subflorescence. Differently, Hall and Hoff (2002) argue that soluble salts do not alter significantly liquid transport properties but that, in contrast, they can induce relevant vapour pressure changes. They maintain that, because soluble salts depress the vapour pressure of the liquid, a slower evaporation rate and, hence, an advancement of the evaporation front towards the outer surface arises during drying. This suggests that the higher tendency of some salt solutions to form subflorescence may arise from a faster evaporation rate rather than from a slower liquid transport.

The higher vapour pressure and, hence, faster evaporation of some salt solutions, for instance sodium sulfate in comparison to sodium chloride, promotes higher supersaturation, thus, higher crystallization pressure, hence, stronger damage (Rodriguez-Navarro and Doehne 1999). This could also account for the higher destructive power of sodium sulfate in laboratory tests.

Temperature-induced crystallization may occur for salts whose solubility is temperature dependent. This is the case of Na_2SO_4 but not of NaCl, which configures another important difference between the two salts. Mirabilite ($Na_2SO_4.10H_2O$) is the stable phase of sodium sulfate below 32.4°C, not only in the air when the RH exceeds that corresponding to the hydration of the salt but also when in contact with a saturated solution (MacMahon *et al.* 1992). Hence, if specimens filled at ambient temperature with a saturated solution are subjected to a temperature drop, mirabilite, whose solubility increases with temperature (Fig. 2.8), may readily crystallise. A similar effect was noticed early on by Brard who reported (Thury 1828), regarding his Na_2SO_4 crystallization test, that "rocks which resist (...) the action of solution saturated when cold, fall apart completely when exposed to the action of solution saturated when hot". Indeed, above 32.4°C thenardite is the equilibrium phase of sodium sulfate. Because the solubility of thenardite is mostly higher than the solubility of mirabilite (Fig. 2.8), when the temperature falls, mirabilite tends to crystallize readily. Doehne *et al.* (2002) indicates that temperature-dependence of salts solubility is one of the features that may enhance crystallization damage in porous materials.

For salts that possess crystal phases with distinct solubility, such as sodium sulfate, crystallization may occur simply when a saturated solution contacts with pre-existing crystals. Doehne *et al.* (2002) indicates this as another feature that results in increased damage to porous building materials. As suggested by Chatterji and Jensen (1989), Flatt (2002) or Tsui *et al.* (2003), sodium sulfate damage in tests involving cycles of impregnation (at ambient temperature) and drying (commonly in an oven) occurs during the impregnation stage due to such an effect. This is, for example, the case of ASTM (2005) test for aggregates or CEN (1999b) test for natural stone. In these tests, very concentrated or saturated solutions are used, respectively. At the end of drying, which takes place at a high temperature in both cases, the porous material contains thenardite because this is the equilibrium phase in the air above 32.4°C. Yet, below 32.4°C the equilibrium phase in contact with saturated sodium sulfate solution is mirabilite. Hence, when thenardite comes into contact with the solution introduced at a new impregnation, a solution highly supersaturated with respect to mirabilite results and crystallization may readily occur.

Salt crystallization growth patterns

By means of experiments on sodium sulfate and sodium chloride, Rodriguez-Navarro and Doehne discovered that some crystals (mirabilite and thenardite) tend to precipitate inside the solution, hence, making the occurrence of subflorescence more likely, while others (halite) tend to precipitate at the air-solution interface. Thus, the formation of efflorescence is more probable for sodium chloride, while for sodium sulfate subflorescence can more easily occur.

Furthermore, the precipitation and growth rate of different crystals may vary under the same test conditions. In the experiments by Rodriguez-Navarro and Doehne, halite showed a slow rate of precipitation and growth. The slower the growth rate, the lower the crystallization pressure. This could also account for a low damage capability of sodium chloride.

Some crystalline habits seem to be associated with more severe damage in porous materials. But all soluble salts can cause severe damage to porous building materials regardless of their equilibrium habits, as stated by Arnold and Zehnder (1985a, 1985b). These authors observed that habits different from the equilibrium ones commonly appeared on walls and argued that crystal morphology is mostly governed by the humidity of the substrate (Zehnder and Arnold 1989). According to them, equilibrium crystal forms are rare in efflorescence, nonetheless they may be found in bristly efflorescence or crusts on very wet substrates. In contrast, whiskers (needle or hair-like crystals) are a common habit for different salts in efflorescence growing on substrates with lower moisture content. Chatterji (2005) also stresses the importance of crystal habits. This author argues that needle-like habits and fast crystal growth are particularly damaging and partially interrelated features. He sustains that fast crystal growth from a highly supersaturated solution gives rise to needle-like habits, fast growth at low supersaturation originates plate-like habits and slow growth from any solution yields equant crystals.

Environmental conditions

Fast evaporation can induce the growth of subflorescence, rather than efflorescence, because the drying front tends to be located deeper in the material. Further, fast evaporation induces higher supersaturation, hence, higher crystallization pressure. The work of Chatterji (2005) suggests that a faster evaporation (hence faster crystal growth and higher supersaturation) gives rise to more damaging habits, namely, needle-like crystals. It is not completely clear to what extent this work contradicts that of Arnold and Zehnder who observed (1985b) that needle-like efflorescence often arises under slow evaporation conditions.

The environmental conditions can cause direct precipitation of crystal phases able to generate greater crystallization pressures. In the case of sodium sulfate, for instance, a temperature above 32.4°C may induce direct precipitation of thenardite, a phase able to generate greater crystallization pressure than mirabilite for equal supersaturation ratios. It is not clear whether precipitation of the heptahydrate, which can occur in metastable conditions bellow 32.4°C, may cause damage. MacMahon *et al.* (1992) consider its formation extremely unlikely under all except laboratory conditions because crystallization of that metastable form requires a very rapid temperature drop and because it is unstable in the presence of mirabilite. Furthermore, Rijniers (2004) argues that this metastable phase is not able to produce crystallization-induced damage in porous materials because its solubility is independent of the pore size.

Substrate properties, namely, porosity and pore size distribution

Schaffer (1932) observed that salt crystallization caused higher damage to stones with small pores than to those with larger pores. Rodriguez-Navarro and Doehne pointed that salt solutions are drawn from the larger to the smaller pores during drying. Hence, crystallization occurs in the smaller pores, the larger pores acting as solution supply reservoirs.

Yet, Scherer (2004) argues that crystallization may take place both in small and in large pores: "under equilibrium conditions, where the crystal is surrounded by a film of solution, high stresses are expected only in small pores, but when that film is discontinuous (as may occur during drying), high stresses can arise even in large pores". Indeed, salt crystallization in large pores (diameter from 1 to 10 μ m) was observed by Zehnder and Arnold (1989) within laboratory tests on brick and mortar samples.

The material porosity and pore size distribution influence also the decay process at a macroscopic level. Rodriguez-Navarro and Doehne argued that small-pored stones are more susceptible to salt decay, to a large extent, because a different hydric flux balance arises during drying: (i) liquid migration occurs more slowly in smaller than in larger pores; (ii) smaller pores result in a larger surface area for evaporation, hence, in a higher evaporation flux. Subflorescence occurs more easily in small-pored stones because drying is faster and, thus, the drying front is located more deeply. Further, a fast evaporation rate can also lead to higher supersaturations, thus, to stronger crystallization pressures. These flux differences may also influence crystal habits, as explained above.

A faster evaporation rate from smaller-pored than from larger-pored stones was also reported by Rousset-Tournier (2001). Moreover, she observed that, with no air velocity, the evaporation rate from all the stones was always higher than from free water surfaces. This suggested that the faster drying rate of the fine-pored stones was due to a higher effective surface of evaporation arising, at least partially, from a higher curvature of the menisci in the small pores.

The mechanical resistance of the material is also a relevant parameter because physical damage occurs when the crystallization pressure exceeds this resistance. Indeed, the stronger the material the less damage is the crystallization process able to cause.

2.2.7 - Working principles of plasters and renders

There are important differences between plasters/renders and other building materials as regards salt crystallization. Firstly, plasters and renders are replaceable. Secondly, they often have the role, which superimposes over their own preservation, of protecting the masonry or adjacent elements from salt decay. Therefore, salt crystallization on or in plasters/renders is not in itself a negative feature.

Failure of plasters and renders occurs when they divert salt solutions to adjacent elements, introduce harmful salts into the masonry or present too low durability. But above all, plasters and renders fail when they do not behave in accordance with their working principle. The working principle of a plaster or render concerns the depth(s) at which salts are expected to accumulate. Therefore, it conditions major use requirements such as the absence of damage at the surface or in the masonry.

There are four main working principles (Wijffels *et al.* 1997, Vegès-Belmin *et al.* 2005, Rodrigues *et al.* 2005) which are expressed in Fig. 2.10 as concern evaporative crystallization processes:

- a) Salt-transporting plasters and renders transport liquid solutions towards the outer surface of walls. Therefore, efflorescence may easily form. This behaviour is expected, for example, from traditional air lime plasters and renders.
- b) Salt-accumulating plasters and renders allow penetration of liquid solutions from the substrate but prevent these solutions from reaching the outer surface. Thus, salt accumulates in the pore system of the plaster/render. Many industrial plasters and renders are formulated on the basis of this working principle. They are particularly interesting when surface damage is undesirable such as in houses, libraries, etc.
- c) Salt-blocking plasters and renders allow vapour but not liquid transport. Therefore, evaporation and crystallization take place at the plaster/substrate interface or in the masonry. This sort of behaviour is expected, for example, from fully hydrophobic materials.
- d) Moisture-sealing plasters and renders do not allow either liquid or vapour transport. Hence evaporative crystallization does not occur. However, two kinds of problems may arise: (i) salt solutions are totally diverted and may damage adjacent elements;
 (ii) if points of escape arise, for instance due to cracking of the plaster/render, salt damage occurs and may progress very quickly, as shown in Fig. 2.10 d).



d) Moisture-sealing: damage sequence due to cracking of the plaster or render

Fig. 2.10 - Representation of the four main working principles of plasters and renders

2.2.8 - Salt crystallization tests

The performance of porous material in relation to salt crystallization and the effects of different salts are currently assessed by means of salt crystallization tests.

Typically, in these tests, the specimens are first contaminated by absorbing the salt solution under a predefined contamination procedure. Afterwards, they are allowed to dry under certain environmental conditions. Normally, several wetting/drying cycles are carried out, though some crystallization tests are performed on specimens in continuous partial immersion. In the last case, absorption and drying occur simultaneously throughout the entire testing period. In crystallization tests, damage is evaluated, either periodically during the test, or at the end, by visual observation or by measuring certain material properties.

Some crystallization tests are the object of official standards, for instance, CEN's test for natural stone (CEN 1999b) or ASTM's test for aggregates (ASTM 2005). Yet, to the best of the author's knowledge, no crystallization tests specific for plasters/renders exist as standard methods.

Salt crystallization testing of porous materials was reviewed by Evans in his fundamental article (1969-1970). More recently, other authors such as Alonso *et al.* (1987) or Goudie and Viles (1997) have also reviewed the theme. But no crystallization tests specific for plasters/renders are mentioned in these review articles, either.

2.3 State of the practice

2.3.1 - Introduction

Salt decay is, in Portugal, commonly called "the cancer of ancient constructions". That shows how serious and hard to deal with practitioners consider this degradation process to be.

To figure out how salt decay problems are handled in practice, end-users from different areas were interviewed (Gonçalves *et al.* 2003). The interviews were carried out under the framework of the COMPASS project simultaneously with others carried out in The Netherlands, France and Spain (TNO 2002). They were mostly aimed at: (i) evaluating the framework in which salt decay problems are approached, namely, the level of knowledge and interaction between the different actors, preliminary investigations carried out and main concerns of practitioners as regards salt damage; (ii) knowing the type of plasters and renders used on salt-loaded walls; (iii) identifying decayed buildings for further selection of case-studies.

A preliminary questionnaire was sent to Portuguese entities and technicians involved in old building conservation: 6 regional delegations of the national authority for monument preservation (IPPAR), 123 local authorities (members of the Portuguese Association of

Municipalities with a Historic Centre), 24 contractors and 1 painter. This questionnaire asked mostly for information on plastered/rendered buildings with salt problems. Answers were received from the 6 regional delegations of IPPAR, 30 municipal authorities and the painter. More than 120 buildings throughout the whole country were indicated. Among the professionals that answered the questionnaire, 2 architects and 1 civil engineer from the national authority, 3 architects and 1 civil engineer from local authorities, as well as the painter, were selected for the interviews. None of the contractors answered the preliminary questionnaire. It was therefore necessary to make direct contacts with some contractors, which allowed the selection of 2 civil engineers and 1 conservator/restorer. In addition, a civil engineer from a consultancy company and an independent consultant (architect), both experts in old building conservation, were selected for the interviews.

A survey was also carried out in the Portuguese market in 2001 (Gonçalves 2002a). It was aimed at identifying industrial plasters/renders or related products available for use on moist salt-loaded walls.

The main conclusions on how salt decay problems are approached in practice are presented in the next section (2.3.2). The types of plasters and renders used in Portugal on salt loaded walls will be focused in section 2.3.3 within a historical perspective. The case-studies will be presented later, in Chapter 6.

2.3.2 - General approach to salt decay problems

The interviews carried out with end-users from different professional areas allowed the following conclusions to be drawn (Gonçalves *et al.* 2003):

- There is a generalized lack of scientific knowledge on salt damage processes among the various actors in the four countries. Only some actors have a reasonably high level of knowledge. In Portugal, only the independent consultant and some technicians from the national authority IPPAR revealed a high level of knowledge. In fact, many endusers in Portugal aim at improving their knowledge on the subject. This was shown by an inquiry performed within a recent seminar on the subject carried out at LNEC (LNEC 2006). Indeed, 78% of the 94 answers received from the 211 participants (professionals from national or local authorities, researchers, academics, contractors, conservatorrestorers, consultants, etc.) indicated the improvement of their theoretical knowledge on salt decay features as one main objective for their attendance at the seminar.
- The responsibility of the various actors (contractors, authorities, suppliers, etc.) is not often clear and the exchange of information between them is frequently insufficient.
- Preliminary inspections or analyses by independent experts are rarely carried out in Portugal, France and The Netherlands.
- In Portugal, end-users in general show more concern about surface than in-depth damage.

 Portuguese end-users complain that salt damage is often aggravated after restoration interventions. This general complaint seems to refer to an increase in surface damage, namely, enlargement of the affected area or more intense dampness, efflorescence or material damage. In many cases the main objective of these interventions was precisely to solve salt damage problems.

The fundamental question that arises from these interviews concerns the factors that can contribute to the worsening of salt decay problems after restoration interventions. One straightforward hypothesis is the type of plasters and renders used, namely, cement-based materials that, as seen in section 2.3.2, can give rise to carbonate or sulfate salts. Decay by these types of salt was identified in many old buildings (Arnold 1981, for instance) but its relevance in Portugal is mostly unknown.

A second important question concerns a possibly wrong approach to salt decay problems. Indeed, the present state-of-the-art does not yet allow a full understanding of many salt decay features. Yet, it is possible that the lack of preliminary investigations, the insufficient background knowledge of many professionals, the unclear responsibilities or the poor interaction between the involved actors impede solving even the simpler problems.

2.3.3 - Main types of plasters and renders used on salt-loaded walls

Air lime plasters and renders were used in ancient times to prevent masonry deterioration, to improve the habitability conditions in buildings or for aesthetical reasons. Under current working conditions, air lime plaster and renders required assiduous (typically annual) maintenance and total replacement in periods of a few years. On salt-loaded walls, even more intense maintenance was most likely needed. Firstly, because lime-based materials have low mechanical strength. Secondly, because lime plasters and renders behave typically as salt-transporting systems and, hence, may show significant surface damage.

Sometimes, additives such as animal or vegetal fats (Rojas 1987, Gonçalves 2002b) were used, aimed at improving the performance of lime-based materials in relation to salt crystallization. It seems reasonable to suppose that hydrophobic (salt-blocking) or else moisture-sealing plasters and renders that postponed the occurrence of surface damage would be obtained. Other techniques existed also to fight against salt damage such as that reported to the author of this thesis by a contractor from Évora (Gonçalves 2002b): the plaster or render was applied over a cotton sheet previously laid on the masonry. The tissue probably created a thin air layer capable of impeding liquid transport from the masonry to the plaster/render. It is possible that the system would therefore work as a salt-blocking system. But there is no scientific evidence on the effective performance and durability of these ancient techniques which have an essentially regional and artisanal character.

After the industrial revolution, Portland cement was eventually introduced as the main binder of plasters and renders. But, despite advantages such as higher mechanical strength and quick hardening even under high RH conditions, cement-based materials often presented low performance on moist and salt-loaded walls. Indeed, worsening of dampness and saltinduced surface damage or detachment from the substrate is regularly observed with cement-based plasters and renders. Further, it was found that cement based materials could give rise to salt damage mostly caused by carbonate salts originating from the (high) alkali content in these materials.

Because of the limitations of lime-based materials and poor performance of cement-based materials, industrial plasters and renders specific for salt-loaded walls have been (and are being) developed in modern times. They are generally dry factory-mixed mortars and are aimed at meeting more demanding requirements of durability and performance.

Industrial salt-blocking plasters/renders based on hydrophobic additives or moisture-sealing plasters/renders based on hydraulic-plus-polymeric binders are common.

The so-called "sanierputzsysteme" (restoration plaster/render systems) were first developed in Germany. WTA established the requirements for such systems in technical specification WTA 1-85, later replaced by WTA 2-2-91 (WTA 1992) and WTA E-2-6-99/D (WTA 2000). WTA systems are composed of dry factory-made mortars that include: (i) hydraulic binders that allow quick hardening and provide good mechanical resistance; (ii) hydrophobic additives that reduce liquid transport; (iii) air-entraining additives or lightweight aggregates that enhance vapour transport. The main purpose of WTA systems is to keep moisture and salts away from the surface and to present good resistance to salt crystallization. They may, therefore, work as salt-accumulating or salt-blocking systems.

More recently, industrial salt-transporting plasters/renders have also been developed to overcome some limitations of salt-accumulating plasters, namely (Hilbert *et al.* 1992, Meier 1992, Hilbert 1995, Goretzki and Karolewski 1995, Hafezi and Figgemeier 1999, Holmström 2000): (i) risk that salts crystallize in the substrate and thereby damage it; (ii) increase in the maximum height of capillary rise in walls due to a reduced moisture transport capability; (iii) uncertain salt-accumulation capability; (iv) progressive decrease in the vapour permeability of the system due to accumulation of salt; (v) poor compatibility with weak substrates. These industrial salt-transporting plasters/renders are aimed at reproducing the moisture-transport and salt-accumulation behaviour of traditional lime-based plasters and renders, while presenting a higher resistance to salt crystallization. They promote the quick transport of liquid solutions towards the outer surface and, hence, a fast drying rate, as it will be explained along this thesis. Yet, dampness and efflorescence are likely to occur with these systems and, therefore, periodic cleaning is necessary.

Recently, European standard EN 998-1 (CEN 2003) framed plasters and renders for salt-loaded walls by defining the concept of "renovation mortar" as being a "designed rendering/plastering mortar used on moist masonry walls containing water soluble salts" that "has a high porosity and vapour permeability and reduced capillary action". The standard does not state which kind of accumulation behaviour is expected from plasters and renders made from these renovation mortars. Nevertheless, the similarity (Table 2.1) between the moisture transport properties in EN 998-1 and those in WTA 2-2-91 (WTA 1992) suggests that they refer to the same type of plaster/render. Yet, it is not clear why CEN requirements for capillary absorption and vapour permeability are only applicable to mortars intended to be used in external elements.

Test parameter	CEN renovation mortars (CEN 2003)	WTA sanierputz (WTA 1992)
Capillary water absorption	Mortars to be used on external elements: ≥ 0.3 kg/m ² after 24 h Test according to EN 1015-18 (CEN 2002)	> 0.3 kg/m ² after 24 h Test according to DIN 52617 (DIN 1987)
Water penetration	≤ 5 mm After the capillary absorption test	< 5 mm After the capillary absorption test
Water vapour diffusion resistance	Mortars to be used on external elements: $\mu \le 15$ Vapour diffusion resistance μ measured according to EN 1015-19 (CEN 1998b)	μ < 12 Vapour diffusion resistance μ measured according to DIN 52615 (DIN 1973)

Table 2.1 – Requirements for moisture transport properties
of CEN renovation mortars and WTA sanierputz

Different kinds of industrial plasters/renders for salt-loaded walls were identified on the Portuguese market (Gonçalves 2002a). Some claim, more or less clearly, to be moisture-sealing, salt-blocking or, although these were mostly introduced in Portugal only around 2000, salt-accumulating. No industrial salt-transporting products were found. Other products that, often with no clearly stated working principle, claim to be adequate for avoiding salt damage were also identified: hydrophobic or moisture-sealing (synthetic-polymer or bituminous) primers, hydrophobic or air-entraining additives for site-made mortars or surface hydrophobic treatments.

As regards the type of plasters and renders used in practice, the interviews carried out (Gonçalves *et al.* 2003) indicated that:

- In Portugal and Spain, traditional plasters and renders, sometimes with hydrophobic or sealing additives, are currently used on moist salt-loaded walls. In the Netherlands and France, it is mostly salt-accumulating or salt-transporting industrial plasters and renders that are used particularly on walls with high moisture or salt content.
- In the four countries, plasters and renders are chosen primarily on the basis of the technicians' own practical experience. In Portugal, end-users with a lower level of knowledge, for instance, professionals from local authorities, are more likely to use plasters/renders with which they do not have experience, namely industrial products. The cost is in general a second order factor, although it seems to be particularly relevant for contractors.
- Several end-users in Portugal and in The Netherlands think that supplier companies do not provide totally independent advice.
- In Portugal and France, the written documentation for industrial products is often unclear or is not in accordance with the stated working principle. For instance, some products are claimed to work as salt-accumulating plasters/renders and, yet, include a strongly hydrophobic first layer.

These interviews revealed a general distrust of the effectiveness of most plasters and renders on moist salt-loaded walls. This is a consequence of recurrent failures experienced with both traditional and non-traditional systems. Accordingly, in Portugal where end-users are more concerned about surface than in-depth damage, hiding the problem was systematically pointed to as the most effective solution. This is normally achieved by building a secondary wall in front of the damaged one. Two end-users (national authority and contractor) also indicated the use of draining geocomposites (a draining nucleus covered with two geotextiles, one on each side of the nucleus) close to the masonry. The lime or lime/cement plaster/render is applied over the geocomposite. An armour of distended metal (fastened to the masonry) or fibreglass (fastened to the geocomposite) is previously placed, to hold the mortar.

The SALTEX dessolidarized plastering system (Holmström 2000) is a somewhat similar solution. It was developed by the Royal Institute of Technology in Stockholm and is composed of: (i) an air layer of 12 to 25 mm that is typically located between the plaster and the masonry; (ii) vapour permeable membrane whose properties are defined on a case-by-case basis according to the type of salts and moisture content in the wall; (iii) one or several traditional mortar layers that may incorporate a metal mesh mechanically fastened to the masonry. Apart from "hiding the problem", this system is aimed at preventing salt crystallization. Although the principle is not totally clear, this purpose is apparently achieved with a vapour permeable membrane by which an RH above the RH_{eq} of the contaminant salt(s) is created in the air layer next to the masonry.

Secondary walls or intermediate air layers are not always feasible because they involve higher costs, especially when compared to traditional plasters/renders, or a sometimes relevant reduction of room area. Also, thick surface layers are frequently not compatible with window frames, doorframes or artistic elements in old buildings. Hence it is not often possible to use them on these buildings when the original aesthetics is to be maintained. Further, masonry preservation is sometimes a real, although not generalized, concern. Particularly in the cases of valuable monuments and end-users with a high level of knowledge, plasters and renders that can effectively work as sacrificial layers are normally sought.

Chapter 3 – General laboratory techniques

3.1 Introduction

Efficient laboratory techniques, for control of the environmental conditions and for evaluation of the salt content in building materials, were required to perform the experimental work in this thesis. Two simple and inexpensive methods were considered for each of these purposes: (i) the use of aqueous solutions for generating constant RH; (ii) the hygroscopic moisture content (HMC) method for assessment of the salt content. However, a question arose as to whether these techniques were fully adequate for testing salt-loaded building materials.

In fact, both techniques required deeper investigation because:

- (i) The use of aqueous solutions for humidity control in the presence of salt-loaded materials had not been, to the best of the author's knowledge, discussed in any publication. In particular, ASTM standard E104-02 (ASTM 2002), which is of wide scope, does not refer that specific kind of material.
- (ii) The HMC method revealed, in recent laboratory research (Lubelli *et al.* 2004), good results as a method for determination of the salt content in porous building materials. However, a systematic analysis of its possibilities and limitations was still necessary, particularly for the testing conditions used in the experiments performed in this thesis.

Efficient experimental methods of general use, as these ones, are an added value for laboratories working on building conservation. Since salt-loaded materials are becoming more and more current in these laboratories, the present study was of broader interest than just the experiments carried out in this thesis.

Aiming at defining adequate guidelines for practice, research work was therefore carried out with the objective of assessing the possibilities and limitations of the two methods. In this chapter, that work is presented. Background information on hygroscopicity is introduced in section 3.2. The method of relative humidity control by aqueous solutions is analysed by means of the experimental research presented in section 3.3. The HMC method is evaluated by means of the theoretical and of the experimental research presented in sections 3.4 and 3.5, respectively. The overall conclusions on the two methods are presented in section 3.6.

3.2 Fundamentals

3.2.1 - Relative equilibrium humidity of aqueous solutions

Molecules in liquids or gases are in constant movement. At a given temperature, liquid substances are continuously evaporating because a number of their molecules possess enough kinetic energy to get away from the liquid surface. At the same time, condensation occurs constantly because some of the evaporated gaseous molecules collide with the liquid surface and are held again by the intermolecular forces in the liquid. In closed systems, a state of dynamic equilibrium, where evaporation is compensated by condensation, is eventually reached between the liquid and its environment.

The pressure exerted under these equilibrium conditions by the (evaporated) gas phase is the "equilibrium vapour pressure" or simply "vapour pressure" of the substance. Vapour pressure increases with increasing temperature because more molecules have enough kinetic energy to leave the liquid. If the liquid is water or an aqueous salt solution, that vapour pressure is the so-called "water vapour pressure" of water or of the salt solution at the given temperature.

The relative humidity (RH) refers to the concentration of water vapour in the air. It corresponds to the ratio, expressed as a percentage, of the actual water vapour pressure in the air to the water vapour pressure above a flat surface of water at the same temperature. In equilibrium conditions, the RH above a liquid is the "relative equilibrium humidity" (RH_{eq}) of the liquid. Thus, the RH_{eq} of a flat surface of pure water is 100% at any temperature. The RH_{eq} above an aqueous salt solution is lower because salt ions replace some water molecules at the liquid surface. Therefore, fewer water molecules are available for evaporation. Accordingly, the evaporation rate, the water vapour pressure and the RH_{eq} of aqueous salt solutions are lower than those of pure water. The more dilute a solution the higher its RH_{eq} at the same temperature. The RH_{eq} of an infinitely dilute solution is 100%.

In Equilibrium Thermodynamics, the RH_{eq} of a solution is expressed as a fraction instead of as a percentage and is called "water activity" (a_w). Eq. 3.1 relates the water activity a_w of a solution to another relevant thermodynamic parameter, the molal osmotic coefficient which expresses the deviation of the solution from ideality (Robinson and Stokes 2002).

$$\ln a_w \equiv -\phi M_w .v.m \tag{Eq. 3.1}$$

In this equation, ϕ (dimensionless) is the osmotic coefficient, M_w (kg/mol) the molar mass of pure water, m (mol/kg) the molality of the solution and ν the stoichiometric parameter (dimensionless) which corresponds to the number of moles of ions produced when a mole of solute molecules is dissociated (Blandamer *et al.* 2005, for instance).

For an ideal solution, ϕ is unitary. The higher the concentration of a salt solution the lower its osmotic coefficient ϕ . Both a_w and ϕ are experimental parameters that can be obtained in published tables of thermodynamic data such as those in Robinson and Stokes (2002) or Lobo (1989). This sort of tables presents a_w or ϕ for different concentrations of a given salt solution at a certain temperature.

3.2.2 - Control of the relative humidity by means of aqueous solutions

The previous considerations show how pure water and salt solutions can influence the RH of the environment. In closed systems, they force the air RH to match the value of their RH_{eq} at the corresponding temperature. This is the basis of the methods of generating constant RH in relatively small containers by means of aqueous solutions, such as those of ASTM E104-02 (ASTM 2002) or ISO 483:2005 (ISO 2005), the latter being specific for plastics. The containers are placed in rooms at a constant temperature. Generally, saturated salt solutions with an excess of salt are used. A wide range of RH can be obtained from different salts: for instance, at 20°C, a saturated caesium-fluoride solution generates 3.8% RH while a saturated potassium-sulfate solution generates 97.6% RH (ASTM 2002). Also pure water is sometimes used, though mostly for research studies, when an RH around 100% is required.

3.2.3 - Hygroscopic behaviour of soluble salts

Soluble salts are hygroscopic which means that, because they have a strong affinity for moisture, they readily attract moisture from the air under certain environmental conditions. Further, they are deliquescent, that is, they eventually dissolve in this hygroscopic water and form liquid solutions.

At a given temperature, both the occurrence of dissolution and the equilibrium concentration of the resulting solution depend on the actual RH of the air and on the relative equilibrium humidity of the salt (RH_{eq}^{sat}). The RH_{eq}^{sat} corresponds to the RH_{eq} of a saturated solution. Three equilibrium situations are possible for non-hydratable salts as NaCl (Fig. 3.1):

- at RH < RH_{eq}^{sat} the salt attracts no moisture from the air;
- at RH = RH_{eq}^{sat} the salt attracts and eventually holds an amount of moisture Q = Q_{eq}, forming a saturated solution;
- at RH > RH_{eq}^{sat} the salt attracts and ultimately holds a given amount of moisture Q > Q_{eq}, forming an unsaturated solution; Q increases exponentially with the air RH; at an RH of 100% the salt solution tends to infinite dilution.



Fig. 3.1 - Hygroscopic moisture held by a non-hydratable salt at a certain temperature. Price and Brimblecombe (1994) present a similar graph for a 1g NaCl sample. For hydratable salts there may be other values of the air RH, below the RH_{eq}^{sat} of the salt, at which the salt attracts moisture from the air and undergoes the corresponding phase change. For instance, sodium sulfate can form three distinct crystalline phases, as explained in section 2.2.4 and depicted in Fig. 2.9: the anhydrous thenardite (Na₂SO₄), the decahydrate mirabilite (Na₂SO₄.10H₂O) and the metastable heptahydrate (Na₂SO₄.7H₂O).

When soluble salts are present in or on porous building materials, either as crystals or salt solutions, their hygroscopic properties may cause or worsen moistening of the material even in the absence of other sources of moisture. Indeed, if the air RH is higher than the RH_{eq}^{sat} of the salt, salt crystals attract moisture from the air and dissolve. Further, while the air RH is higher than the RH_{eq} of the salt solution, either this solution was newly formed by deliquescence of pre-existing crystals or was previously present in the material, a negative vapour pressure gradient exists between the salt-loaded material and the surrounding air. Hence, water vapour diffuses from the environment to the surface and, thus, the water content in the material increases. In contrast, when the environmental RH is lower than the RH_{eq} of the salt solution, the vapour pressure gradient is positive. Therefore water vapour diffuses in the opposite direction, that is to say, from the material to the environment. Hence, the salt-loaded material dries.

3.3 Use of aqueous solutions for control of the relative humidity

3.3.1 - Introduction

Methods of controlling the RH by means of aqueous salt solutions are widely used for storing or testing building materials. Relatively small containers where the RH is controlled by chosen solutions are utilized, placed in rooms at a constant temperature. Indeed, stable and accurate RH is much more difficult to obtain in laboratory rooms than temperature. The low cost and simplicity of such methods of RH control make them a good alternative to climatic chambers. These chambers are large, expensive and energy consuming equipments whose availability is normally restricted and needs to be rationalized.

In the case of salt-loaded materials, accurately controlled environmental conditions are often indispensable because the behaviour of soluble salts depends on the air RH (and temperature). Yet, as far as the author knows, the correctness of using aqueous solutions for generating a constant RH when the stored materials contain soluble salts had never been evaluated before this thesis.

In such methods of RH control, selected salt solutions or even pure water are used to generate a constant RH (identical to their RH_{eq}). When wet materials that contain soluble salts are stored in the containers, secondary salt solutions are also present, which supposedly may influence the environmental conditions as well. Yet, the only thing currently taken into consideration is that wet materials can affect the environmental conditions, particularly in the case of low RH and high temperature, because water vapour can be released and because evaporation consumes heat.

Accordingly, no major problems are expected from dry materials. However, soluble salts are deliquescent. Consequently, secondary salt solutions can form also on salt-loaded materials even when they are initially dry.

Research was, therefore, undertaken (Gonçalves and Abreu 2007) with the main purpose of verifying whether the use of aqueous solutions allows stable control of the RH in the case of salt-loaded materials. Its specific objectives were:

- To test, by means of adequate experiments, the null hypothesis that stable environmental conditions are always obtained with dry materials even if they contain soluble salts;
- To examine the case of wet materials.

This section presents and discusses the research carried out. Two experiments are described where NaCl samples were kept over water in section 3.3.2. Then, the results are presented in section 3.3.3 and discussed in section 3.3.4.

3.3.2 - Materials and methods

The two experiments were performed by placing NaCl samples ($RH_{eq}^{sat} \approx 75.6\%$ at 20°C) of 1g and 5g, respectively, inside a closed container over water ($RH_{eq} = 100\%$). The RH in the container, a small glass desiccator cabinet, was monitored through periodical observations by means of a thermo-hygrometer. This experimental set-up can be seen in Fig. 3.2. The two experiments were carried out in similar conditions, one after the other, in a conditioned room at 20°C and 50% RH.

3.3.3 - Results

The results of the experiments are shown in Figs. 3.3 to 3.5. It is seen that the RH inside the container decreased rapidly from the early moments of either experiment. In the case of the 1g sample, this happened after an initial RH increase. Minimum RH of 91% or 85% was attained after 2 and 5 days for the 1g and the 5g sample, respectively. Afterwards, the RH started to rise again. In the case of the 5g sample, a secondary RH drop is clearly visible between the 8th and the 12th day: a second RH minimum of around 85% was reached on the 12th day. In the case of the 1g sample, only a slight irregularity was detected between the 3rd and the 4th day.

A sharp fall in the temperature in the container was detected in both experiments when the RH dropped (Figs. 3.4 and 3.5). In the case of the 5g sample, the temperature fall was simultaneous to the second RH drop. However, since in this experiment the initial temperature values were not recorded, it is not known if there had been a previous temperature fall during the first RH drop.

For the 1g sample, equilibrium of the RH inside the container was attained at the expected value of 100% after 16 days of testing. In the case of the 5g sample, RH of around 99% was recorded, during four consecutive days, only after 56 days of testing.

The amount of water in the samples was visually observed to increase during the experiments.



Fig. 3.2 - Experimental set-up



Fig. 3.3 - Evolution of the RH in the container where NaCl samples were stored over water



Fig. 3.4 - 1g NaCl sample: RH and T in the container

Fig. 3.5 - 5g NaCl sample: RH and T in the container

3.3.4 - Discussion

The results presented above indicate that aqueous solutions may not allow an adequate control of the environmental conditions when salt-loaded materials are stored in the container. Indeed, secondary solutions with a distinct RH_{eq} may influence the air RH. Secondary solutions exist on wet materials or may form, as seen in the present experiments, on dry materials when the air RH is above the RH_{eq}^{sat} of the contaminant salt. Moreover, enthalpy changes may cause disturbance in the temperature. These enthalpy changes can arise from dissolution, as in these experiments, or crystallization processes.

In the present experiments, the NaCl samples were stored at an RH that, under the influence of pure water, tended to rise (towards 100%) immediately after the container was closed. That is clearly seen in the case of the 1g sample (Fig. 3.4).

Although disturbed by the opening of the container, the RH in the interior was, at beginning of both experiments, well above the RH_{eq}^{sat} of NaCl (75.5%), as seen in Fig. 3.3. Therefore, deliquescence of the salt took place and NaCl solutions were formed.

The initial RH drops most likely resulted from the influence of those NaCl solutions whose early RH_{eq} was that of a saturated NaCl solution (75.5%). Indeed, the actual RH in the container reached a lower value when more NaCl was present: 85% RH for the 5g sample and 91% RH for the 1g sample.

The sharp temperature falls (Figs. 3.4 and 3.5) were most likely a result of enthalpy changes due to NaCl dissolution being endothermic. It is interesting to note that a mere 1g of NaCl was able to lower the temperature in the container by 4°C to 5°C during around 24 hours.

The further rise of the actual RH in the container observed in both experiments arose presumably from a progressively lower influence of the NaCl solutions, as they became gradually more dilute and, hence, their $\rm RH_{eq}$ increased. This is consistent with the RH increasing faster in the case of the 1g sample that corresponds to a lower amount of salt solution.

Temperature drops cause an increase in RH of the air. This is probably the case of the small RH increase observed for the 5g sample between the 6th and the 10th day. Such an effect is not clear for the 1g sample, though it may very well have happened at the same time than the initial RH increase

Equilibrium was eventually attained, perhaps not fully in the case of the 5g sample, but only after several days. The higher the amount of NaCl in the sample the longer the period before constant RH was registered: 16 and 56 days for the 1g sample and the 5g sample, respectively.

These results show, therefore, that when the RH is controlled inside small containers by means of water, it can be significantly influenced by the presence of soluble salts stored in those containers. This conclusion can be generalized to any conditioning solution with RH_{eq} higher than the RH_{eq}^{sat} of the contaminant salt because pure water corresponds to a solution with $RH_{eq} = 100\%$.

In fact, if the air RH is above the RH_{eq}^{sat} of the contaminant salt, that salt eventually dissolves. Disturbance of the environmental conditions may arise firstly from the dissolution process itself (for example, NaCl dissolution is endothermic). Then, a dynamic process takes place where the air RH is drawn down by the contaminant solution towards a value that gradually increases as a result of the dilution, and corresponding increase of the RH_{eq}, of the contaminant solution.

Equilibrium is reached only after a certain period of time. That period is probably longer when: (i) the materials contain a high amount of salt; (ii) the RH_{eq}^{sat} of the contaminant and conditioning salts are very different, hence, the contaminant solution tends to a high dilution.

In practice, most problems arise perhaps with dry salt-loaded materials that are stored, as described above, at an RH higher than the RH_{eq}^{sat} of the contaminant salts. But problems can be foreseen also with wet materials, for instance, with samples collected from damp

salt-loaded walls, as well as within wet/dry-cycle tests on salt-loaded materials when RH control by salt solutions is used somewhere along the test. The following situations might theoretically occur:

- a) If the RH is lower than the RH_{eq}^{sat} of the contaminant salt: (i) while the contaminant salt is in solution the RH is drawn to a higher value; (ii) the contaminant salt eventually crystallizes which may induce additional disturbance of the environmental conditions due to enthalpy change (NaCl crystallization, for instance, is exothermic).
- b) If the RH is higher than the RH_{eq}^{sat} of the contaminant salt: (i) the RH is drawn towards lower or higher values when it is initially higher or lower, respectively, than the RH_{eq} of the contaminant solution; (ii) RH deviation decreases progressively as the contaminant solution attracts or loses moisture and ultimately becomes more dilute or more concentrated, respectively; (iii) temperature disturbance could in principle arise from the dilution or concentration process, though its significance is probably low.

Relevance and duration of the environmental disturbance is expected to depend on multiple and interrelated factors such as the nature of the involved salts, exposed area of the conditioning solution, volume of the container, salt content in the stored materials, temperature, etc. Further, salt dissolution and crystallization are complex processes where kinetic factors depending on the type of salt, physical characteristics of the loaded material, environmental conditions, etc. are also possibly relevant. Moreover, multiple situations may occur as regards both the nature of contaminant salt, which can even be a mixture of several salts, and the type of loaded materials.

Nevertheless, it is reasonable to assume that RH disturbance will be less significant when: (i) the salt content in the material is lower; (ii) the difference between the RH_{eq} of the conditioning solution and the RH_{eq} of the contaminant solution (or RH_{eq}^{sat} of the contaminant salt, in the case of dry materials) is smaller; (iii) the exposed area of the conditioning solution is higher; (iv) the storage period of the materials in the container is longer.

3.4 Fundamentals, scope and accuracy of the HMC method

3.4.1 - Introduction

The HMC method of evaluating the salt content in porous building materials is expected to be a quick and inexpensive alternative or complementary method to chemical analysis, namely, ion chromatography (IC) or even conductivity-based methods (UNI 2003, for instance).

Indeed, the study of salt-damaged buildings and the evaluation of specimens from laboratory crystallization tests often involve an evaluation of the salt content of large numbers of samples. This was the case, for instance, during the work presented in Chapters 4 and 5 of this thesis. Yet, chemical analyses are time consuming, require specifically trained personnel and, thus, are often too expensive or too slow to be performed systematically on large sets of samples.
The HMC method, by contrast, requires only a small amount of laboratory work of a trivial nature: introducing the samples into and removing them from a climatic chamber where they are subjected to certain environmental conditions until hygroscopic equilibrium is reached, plus two or three weighing sequences. The HMC method permits, therefore, the simultaneous testing of a very large number of samples, from several dozen to as many as several hundred, without the need to carry out any special preparation of those samples (salt extraction or even grinding) and involving only an insignificant investment of time.

The HMC method of salt content evaluation is primarily based on two main facts:

- Soluble salts are generally very hygroscopic. Hence, they are able to attract and hold much higher amounts of moisture from the air than ordinary building materials such as mortars, stone or brick. Hence, the HMC method can be used to detect the presence of salts (BRE 1989).
- ii) In equilibrium under specific environmental conditions the hygroscopic moisture content of a given salt or salt mixture takes a precise value (Price and Brimblecombe 1994, for instance). Thus, the method is expected to allow quantifying the salt content, at least in relative terms among different samples contaminated with the same salt or salt mixture.

A related test method was three decades ago presented by Rodrigues (1976) to estimate the clay content of natural stone, by measuring the HMC after 72 hours of exposure to an atmosphere with controlled RH. But the origins of the HMC method are founded mostly on BRE Digest 245 (1989), which derived from BRE TIL 29 (1977), even though salt content determination is not the objective of this BRE's method. Here, the HMC, measured at room temperature and 75% RH, is used to evaluate to what extent the actual moisture content in samples collected from damp buildings could be due to hygroscopicity rather than to rising damp.

Wijfels *et al.* (1997) used the HMC method, carried out under controlled temperature and RH, for a relative evaluation of the salt content in NaCl-loaded samples collected from specimens used in crystallization tests. More recently, Lubelli *et al.* (2004) reported an extensive set of HMC experiments performed under controlled temperature and RH on brick samples with known contents of NaCl, Na₂SO₄, NaNO₃ or mixtures of the three salts. These experiments tested the use of the HMC method for a quantitative determination of the salt content. Good results were obtained for all salts.

The research presented in this section (Gonçalves and Rodrigues 2006) gives continuation to these works, particularly to that of Lubelli *et al.* The use of a climatic chamber instead of RH control by aqueous solutions, unlike in these previous works, is justified by the results presented in section 3.3. Moreover, it was verified (Vergès-Belmin 2003) that HMC measurements can take a long time (in the order of months) when performed in closed containers with no air movement.

In practice, climatic chambers do not generate absolutely constant T and RH. Therefore, because the HMC of soluble salts depends on the environmental conditions, there is a hypothesis of an uncertainty being associated to the measured HMC values. If the environmental parameters have a strong fluctuation, it may even be impossible to reach an equilibrium condition, i.e., to obtain stable HMC values.

The present research was aimed at clarifying the possibilities and limitations of the HMC method by means of a theoretical study. It had four main objectives:

- systematizing the HMC method;
- investigating its stabilization conditions;
- assessing its accuracy;
- discussing its scope.

This section presents that theoretical research. The method is systematized in section 3.4.2 and its stabilization conditions, accuracy and scope are discussed in section 3.4.3.

3.4.2 - Systematization of the HMC method

3.4.2.1 - Measurement of the HMC

In order to measure their HMC, the samples are first oven-dried and then weighed. Afterwards, they are stored at the selected environmental conditions for a period sufficiently long for hygroscopic equilibrium to be achieved, that is, until their mass remains constant over time.

Where m_{dry} (g) is the dry mass of a given sample, m_s (g) its mass at hygroscopic equilibrium and $Q=m_s-m_{dry}$ the quantity of hygroscopic moisture in the sample, the HMC (%) of that sample is given by Eq. 3.2:

$$HMC = 100.\frac{Q}{m_{dry}}$$
(Eq. 3.2)

3.4.2.2 - Determination of the salt content

Soluble salts hold, in equilibrium conditions under given environmental conditions, a precise amount of hygroscopic moisture. The same happens to common building materials such as mortar, stone or brick. Therefore, for samples composed of a single material, the amount of hygroscopic moisture Q is directly proportional to the dry mass m_{dry} of the sample, that is, Q/m_{dry} is constant. Hence, at given temperature and RH, the HMC is a constant characteristic of each material, either a salt or a building material, as emerges from Eq. 3.2. This is shown in Fig. 3.6.

If the sample is composed of a base-material loaded with salt, a different situation occurs. The dry mass of the sample is the sum of the dry masses of the two materials: $m_{dry} = m_{salt} + m_{mat}$. The salt content in the sample is $x=100.m_{salt}/m_{dry}$ (%). Disregarding any chemical interaction between the two materials, the amount Q of hygroscopic moisture in the sample can be assumed as the sum of the hygroscopic moisture in either material: $Q=Q_{salt}+Q_{mat}$. Therefore, from Eq. 3.2, the HMC (%) of the composite sample is expressed by:

$$HMC = (a-b)x + 100b$$

In this equation, a and b (g/g) represent the weight amount (g) of hygroscopic moisture in samples with unitary mass (1g) of the salt and of the base-material, respectively. This implies that the HMC of a salt-loaded sample is a linear increasing function of its salt content, as depicted in Fig. 3.7.



Fig. 3.6 - HMC of single material samples



In practice, parameter *b* in Eq. 3.3 and Fig. 3.7 can be disregarded because the hygroscopicity of building materials is not normally significant when compared to that of soluble salts. Hence, *b* can be assumed as zero and the HMC linear function in Fig. 3.7 assumed to cross the origin instead of point B. Therefore, where x (%) is the salt content in the sample and HMC_{salt} the HMC (%) of the salt which is a constant, as seen in Fig. 3.6, the HMC of the salt-loaded sample can be assumed as:

$$HMC \approx ax \quad \iff \quad HMC \approx \frac{HMC_{salt}}{100}x$$
 (Eq. 3.4)

Eq. 3.4 shows that the HMC of a salt-loaded sample depends essentially on: (i) the nature of the salt; (ii) the salt content in the sample.

Parameter a may be calculated by one of the following methods:

i) By means of tables of thermodynamic data of the salt at the given temperature. Indeed, the actual RH in the chamber corresponds to the water activity of the salt solution that forms at hygroscopic equilibrium. If that actual RH is assumed to correspond to the set-point RH, the molality of this solution can be obtained, by interpolation, from the water activity values in such tables. Both water activity/molality or, by means of Robinson-Stokes equation (Eq. 3.1), osmotic coefficient/molality tables can be used. Where m (mol/kg) is the molality of the solution and M_{salt} (g/mol) the molar mass of the salt, parameter a is then given by:

$$a = \frac{10^3}{m.M_{salt}}$$
(Eq. 3.5)

ii) By performing HMC tests on control-samples with known salt content. This allows obtaining an experimental line that correlates the HMC to the salt content. That correlation line corresponds to Eq. 3.4 when only control-samples of the pure salt are used. If control-samples of the base-material are also used, the correlation line corresponds to Eq. 3.3.

3.4.3 Discussion

3.4.3.1 - Accuracy of HMC measurements

For soluble salts, the slope of the curve HMC-versus-RH increases sharply as the values of the RH rise above the RH_{eq}^{sat} . This was qualitatively shown in Fig. 3.1. Fig. 3.8 illustrates that sort of behaviour for six common soluble salts within an RH range of 93% to 97% at 25°C.



Fig. 3.8 – HMC of six soluble salts as a function of the RH. Obtained from equilibrium thermodynamic data for NaCl, KCl, NaNO₃ and KNO₃ (Robinson and Stokes 2002), Na₂CO₃ (Robinson and Macaskill 1979) and K₂CO₃ (Roy *et al.* 1984).

That behaviour implies that: (i) at high RH, small RH deviations may result in significant HMC deviations; (ii) the higher the RH, the larger the HMC variations induced by small fluctuations of the RH. Fig. 3.8 shows that, for instance, a deviation of \pm 1% around 94% RH would cause the HMC of pure NaCl samples to diverge by +19% or -13%, respectively, as a percentage of the HMC at the central RH of 94%. However, a variation of the same magnitude around 96% RH would cause the HMC to vary by +32% or -19%, respectively. Therefore, since climatic chambers do not generate completely accurate environmental conditions, particularly of RH, the measured HMC values can be affected by a considerable uncertainty, especially at high RH.

In this chapter, only RH rather than also temperature deviations will be considered. This simplification is expected to be reasonable because, at high RH, which is the case here, soluble salts are typically more sensitive to RH than to temperature variations. Moreover, in climatic chambers, RH is more easily destabilized than temperature. But, strictly speaking, HMC deviations may also arise from temperature fluctuations.

Consider a salt-loaded sample kept inside a climatic chamber where the temperature is constant but the actual RH may diverge from a minimum value RH^- to a maximum value RH^+ below and above the set-point value RH^e , respectively. As a consequence of this RH variation, the actual HMC of the sample will fall within the interval [HMC⁻(x), HMC⁺(x)]. HMC⁻(x) and HMC⁺(x) are the linear HMC functions (Eq. 3.4) for the maximum and the minimum RH, respectively. That behaviour is depicted in Fig. 3.9 where HMC^e(x) is the linear HMC function for the set-point RH.



Fig. 3.9 - HMC variation due to deviation of the RH

In Fig. 3.9, E^- , E and E^+ (%) represent the HMC of the pure salt at the minimum, the set-point and the maximum RH, respectively. From Eq. 3.4, the amplitude Δ HMC(x) of the interval for the possible HMC values is also an increasing linear function of the salt content x:

$$\Delta HMC = \frac{E^+ - E^-}{100} x \tag{Eq. 3.6}$$

The uncertainty Δx that arises when the salt content x is estimated by means of the actual HMC is also an increasing linear function of x:

$$\Delta x = \frac{100}{E} \Delta HMC = \frac{E^+ - E^-}{E} x$$
(Eq. 3.7)

The significance of Δ HMC or of Δx in relation to HMC^e or x, respectively, is expressed by parameter α (%):

$$\alpha = 100 \frac{\Delta HMC(x)}{HMC^{e}(x)} = 100 \frac{\Delta x}{x}$$
(Eq. 3.8)

Parameter α is a constant, as it elapses when Eq. 3.4, 3.6 and 3.7 or 3.8 are merged:

$$\alpha = 100 \frac{E^+ - E^-}{E}$$
(Eq. 3.9)

Eq. 3.9 shows that α depends essentially on the following factors: (i) type of salt; (ii) set-point values of the environmental parameters; (iii) possible deviation of the environmental parameters from the set-point values, hence, capability of the climatic chamber to generate homogeneous and accurate environmental conditions.

The acceptability of α depends on the specific purposes of the HMC test and, thus, requires a case-by-case evaluation. Consider the example depicted in Fig. 3.10 where the HMC of NaCl-loaded samples is measured in a climatic chamber at 25°C and 95.0±0.2% RH or

95.0±1.0% RH. In the first case, from water activity/molality tables (Robinson and Stokes 2002), E⁺=1196.4%, E=1153.4% and E⁻=1110.4%, thus, $\alpha \approx 7.5\%$. This value is probably acceptable in many practical cases, for instance, when evaluating the deposition of salt across the thickness of specimens from crystallization tests. However, for an RH of 95.0±1.0%, E⁺=1423.0%, E=1153.4% and E⁻=968.8% which gives $\alpha \approx 39.4\%$, hardly an acceptable value in most situations.



Fig. 3.10 – Possible divergence in the HMC of NaCI-loaded samples due to deviation of the RH from the value of 95.0%

An accurate prediction of α for a certain climatic chamber is only possible for salts with known thermodynamic properties. For other salts or salt mixtures, an approximate evaluation of α can, though, be made by means of, for example, the properties of NaCl. Since NaCl is one of the common soluble salts with higher hygroscopic absorption (Fig. 3.8), this evaluation will most likely err on the side of safety.

However, as it emerges from the considerations above, it is probable that most climatic chambers presently available in research laboratories may induce significant error in the HMC measurements.

3.4.3.2 - Mass stabilization

When measuring the HMC, samples should remain under the chosen environmental conditions until their mass is constant over time. Yet, the environmental fluctuations in the chamber can cause variations in the mass of the sample. Therefore, if the environmental fluctuations are too strong, it may be impossible to obtain a stable HMC.

Consider a sample with dry mass m_{dry} (g) and salt content x (%). The mass variation between consecutive weighing *i*-1 and *i* is $\Delta m = m_i - m_{i-1}$. Hence, the mass variation Δm (%) that corresponds to a maximum HMC variation ΔHMC (Eq. 3.6) is given by Eq. 3.10 which derives from Eq. 3.2 and 3.6:

$$\Delta m = \frac{m_{dry}}{10^4} (E^+ - E^-) x \tag{Eq. 3.10}$$

This equation shows that Δm is an increasing function of the salt content *x*. This means that, the higher the salt content in the sample, the more significant are the possible mass variations caused by fluctuation of the environmental parameters.

Therefore, it is extremely difficult to establish a reasonable mass stabilization criterion. Indeed, the usual criterion of considering that equilibrium is achieved when Δm is lower than a certain percentage β of the dry mass m_{dry} (in drying procedures, normally β =0.1 for a 24-hour weighing interval) is not often applicable. This criterion is expressed by Eq. 3.11:

$$\Delta m \le \beta \frac{m_{dry}}{100} \tag{Eq. 3.11}$$

However, since Δm is a function of *x* (Eq. 3.10), for each value of β there is a critical salt content x_{crit} above which mass variations greater than Δm can arise simply due to environmental fluctuation. In that case, the criterion in Eq. 3.11 may never be fulfilled. By replacing Eq. 3.10 in Eq. 3.11, x_{crit} is obtained:

$$x_{crit} = 100 \frac{\beta}{E^+ - E^-}$$
(Eq. 3.12)

The problem is that, in practice, samples with salt content $x>x_{crit}$ can easily occur. For instance, β is taken as 1 (already much higher than the value of 0.1 usual in drying tests). If, for instance, NaCl-loaded samples are tested in a climatic chamber at 95.0±0.2% RH and 25°C, the maximum critical salt content in Eq. 3.12 is given by $x_{crit} = 100*1/(1196.4-1110.4) \approx 1.2\%$ which is an extremely low value. Indeed, in most practical situations, samples with a higher salt content may exist, as it will be seen along this thesis.

Therefore, in practice, it is preferable to qualitatively assess mass stabilization of the samples by direct observation of the mass-versus-time or HMC-versus-time graphs.

3.4.3.3 - Scope of the HMC method

Lubelli *et al.* (2004) concluded that the HMC method may allow quantifying or roughly evaluating the salt content in samples loaded with known or unknown salts, respectively. The present thesis supports, overall, these conclusions.

The scope of the HMC method can be summarized as follows:

- (i) If we are dealing with a salt of well-known thermodynamic properties or if we are able to replicate the salt in the laboratory, the HMC method may be used for a quantitative evaluation of the salt content, as explained in section 3.4.2.2:
 - For salts whose thermodynamic properties are known, correlation parameter *a* in Eq. 3.4 can be obtained from tables of thermodynamic data;
 - For replicable salts, either their thermodynamic properties are or not known, Eq. 3.4 can be directly obtained by using control-samples of the salt. This equation is usually sufficient because the HMC of most base-materials is negligible in comparison to the HMC of soluble salts. If control-samples of the base-material are also used Eq. 3.3 can be obtained.

(ii) If the thermodynamic properties of the salt are not known and that salt is not replicable in the laboratory, since the HMC is proportional to the salt content, the HMC method may still be used, but only for comparing the salt content in different samples. This application rests, however, on the assumption that the type of salt is identical in all the samples. If the type of salt shows a significant variation from sample to sample, the method can still be used but it will only provide more or less rough estimates of the salt content. The homogeneity of the type of salt within a group of samples may be evaluated by performing IC analyses on some selected samples.

Salts of known thermodynamic properties are normally present, for instance, in laboratory samples. On the contrary, samples from buildings are commonly loaded with salts whose thermodynamic behaviour is not known. Sometimes the nature of the salt itself is unknown but what normally happens is that a complex mixture of different salts exists. These complex salt mixtures are not easy to reproduce in the laboratory, adequate thermodynamic data is not normally available for them and the relative proportions of the different salts may show differences from sample to sample.

3.5 HMC experiments on salt-loaded materials

3.5.1 - Introduction

Complementarily to the theoretical research presented in the previous section, the possibilities and limitations of the HMC method were investigated by means of experimental research.

The HMC method is expected to be effective if an RH above the RH_{eq}^{sat} of the contaminant salt is used. In principle, the HMC of the base-materials can be disregarded as it is expected to be insignificant when compared to the HMC of soluble salts. Accordingly, the state of cohesion of the samples is not expected to have a relevant influence on the results.

Nevertheless, stable HMC values can only be obtained if the fluctuation of the environmental parameters is low enough to allow stabilization of the mass of the sample. As discussed in the previous section, there is a strong risk that RH fluctuation in common climatic chambers is not sufficiently low to allow such stabilization. Yet, it is possible that the hygroscopic response of the salts is not immediate and, hence, HMC fluctuation is smoothed in practice. For instance, the shape of the receptacle where the sample is placed is likely to influence the dissolution kinetics.

If stable actual HMC values are eventually obtained, there is still a possibility that they will not correspond to those expected from equilibrium thermodynamic data. Indeed, some soluble salts often present metastable behaviour. That is the case of, for instance, sodium sulfate which was used in several of the experiments in this thesis. Crystallization of the heptahydrate or situations of metastable equilibrium of thenardite have been reported, for example, by Rijniers (2004) or Tsui *et al.* (2003), respectively. HMC values different from the expected ones may also arise due to permanent or erratic deviations of the RH or temperature in the chamber, as discussed in section 3.4. Anyway, regardless of the cause, when the actual HMC deviates from the value that corresponds to the set-point conditions of

temperature and RH, there is an error associated to salt content estimates made on the basis of these HMC values.

A linear correlation HMC/salt content is expected to be obtained by means of samples with different salt content only when: (i) the salt solutions that form by hygroscopic absorption of moisture achieve all their equilibrium concentration; (ii) the environmental conditions are spatially homogenous in the chamber, in the case of several measurements carried out simultaneously; (iii) the environmental conditions are reasonably constant in time, in the case of several series of measurements.

Samples from decayed buildings usually contain a mixture of different salts. The first question that arises is, therefore, whether the HMC of a salt mixture can be calculated by means of the HMC of the different salts. In fact, it may be impossible to directly deduce the properties of salt mixtures, the RH_{eq}^{sat} for instance, from the properties of the individual salts because these interact when in solution, as explained in section 2.2.4. The more concentrated the solution, the stronger that interaction. However, when the HMC is determined at high RH, for instance 95%, dilute salt solutions can form because the RH_{eq}^{sat} of many soluble salts is much lower than that. In this case, it is possible that the HMC of some salt mixtures is similar to the value obtained by weighing the HMC of the individual salts.

A second question concerns the sensitivity of the HMC to variations in the composition of the salt mixture. Indeed, homogenous salt mixtures hardly exist throughout a building or even a wall. Even in the most favourable cases, a certain variation in the relative proportions of the different salts is likely to occur. And, as argued in the previous section, the less constant the type of salt from sample to sample, the less reliable is the HMC method expected to be.

The experimental research presented in this section was aimed at testing the general hypothesis that the HMC method is effective when the RH_{eq}^{sat} of the salts is below the RH of the testing environment. Two base-materials (lime mortar and brick), nine soluble salts (NaCl, KCl, Na₂SO₄, K₂SO₄, NaNO₃, KNO₃, Na₂CO₃, K₂CO₃ and CaSO₄.2H₂O) and four different mixtures of NaCl, Na₂SO₄ and NaNO₃ were used. Most of this research, except the work on salt mixtures, was published as a journal article (Gonçalves *et al.* 2006a).

The main objectives were to assess, for a given climatic chamber and certain environmental conditions:

- the influence of the two base-materials;
- the influence of the state of cohesion of the sample;
- the mass stabilization conditions for two different sample receptacles;
- whether linear correlations HMC/salt content were obtained;
- the accuracy of the measurements, namely as regards: agreement of the actual HMC with the values expected at equilibrium; error associated to salt content estimates;
- for the salt mixtures: agreement of the actual HMC with the value calculated by weighing the HMC of the individual salts; impact on the HMC of varying the relative proportions of the individual salts.

The experiments carried out are described in section 3.5.2 and their results presented in section 3.5.3. These results are afterwards discussed, as regards: influence of the base-materials and state of cohesion (section 3.5.4.1), mass-stabilization (section 3.5.4.2) correlation HMC/salt content (section 3.5.4.3), accuracy of the method (section 3.5.4.4) and behaviour of the salt mixtures (section 3.5.4.6). Sodium sulfate, which behaved differently from the other salts, is more carefully discussed in section 3.5.4.5.

3.5.2 - Materials and methods

Two sets of HMC experiments were carried out, as indicated in Table 3.1, on the following types of samples:

- i) base-materials: red ceramic brick or lime mortar;
- ii) individual salts: NaCl, KCl, Na₂SO₄, K₂SO₄, NaNO₃, KNO₃, Na₂CO₃, K₂CO₃ or CaSO₄.2H₂O; their RH_{eq}^{sat} at 20°C is indicated in Table 3.2;
- iii) each of the base-materials with increasing content of each of the nine salts;
- iv) lime mortar with increasing content of each of four different mixtures of NaCl, Na₂SO₄ and NaNO₃; composition of these mixtures is presented in Table 3.3.

Set	Contaminating salts		Sample type	Dry mass	Receptacle type	Environmental conditions and duration of the test	
1 st	none	i)	lime mortar or brick			0	
	NaCl or Na ₂ SO ₄	iii)	lime mortar or brick with increasing content of either salt	2 g	deep-narrow	- 13 weeks at 20°C / 95% RH	
	none	i)	lime mortar (powder, granular and entire) and brick (only powder)	2 g	deep-narrow		
	NaCl, KCl, Na ₂ SO ₄ , K ₂ SO ₄ , NaNO ₃ ,	ii)	i) nure salt samples	0.5 g	deep-narrow	20°C / 95% RH	
nd		",	pure sait samples		shallow-wide		
2 nd	KNO ₃ , Na ₂ CO ₃ , K ₂ CO ₃ or CaSO ₄ .2H ₂ O		lime mortar with increasing content of each of the nine salts	2 a	doop parrow	 Samples (ii): 64 days Samples (iii) and (iv): during the first 29 days 	
	Four different mixtures of NaCl, Na ₂ SO ₄ and NaNO ₃	iv)	lime mortar with increasing content of each of the four salt mixtures	2 y	deep-narrow		

Table 3.1 - HMC experiments

The lime mortar was obtained from prismatic specimens with volumetric composition of 1:1.5:1.5 (dry hydrated lime LUSICAL : sand from the Tagus river : yellow pit sand from Corroios) prepared at LNEC one year before testing. The brick was obtained from solid red Dutch ceramic bricks. Both the lime mortar and the brick are the same materials used in the drying and crystallization experiments presented in Chapters 4 and 5. The salts were commercial products of "for analysis" quality.

Table 3.2 - Relative equilibrium h	humidity of the	salts
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Salt	RH _{eq} ^{sat} at 20 °C (%)
NaCl	75.47 ⁽¹⁾
KCI	85.11 ⁽¹⁾
NaNO ₃	75.36 ⁽¹⁾
KNO ₃	94.62 ⁽¹⁾
K ₂ SO ₄	97.59 ⁽¹⁾
K ₂ CO ₃	43.16 ⁽¹⁾
Na ₂ SO ₄	95.6* ⁽²⁾
Na ₂ CO ₃	91.6 ⁽³⁾
CaSO ₄ .2H ₂ O	99.6 ⁽⁴⁾

Table 3.3 – Composition of the salt mixtures

Mixture	Relative amount of salt in the mixture (%W)			
Winkton	NaCl	Na_2SO_4	NaNO₃	
A	100/3 100/3		100/3	
В	50	25	25	
С	50 -		50	
D	50	50	-	

* deliquescence RH for $Na_2SO_4.10H_2O$

(1) Greenspan (1977)

(2) Linnow et al. (1989)

(3) Apelblat and Manzurola (2003)

(4) Zehnder (1996)

The samples were prepared in the laboratory after grinding in an agate mortar and then oven drying (at 60°C) both the base-materials and the salts. The composite samples were obtained by simply mixing the powders in the testing receptacles. The entire or granular lime mortar samples were obtained by breaking or disintegrating the mortar prisms, respectively.

Brick or mortar samples with salt contents up to 10% were tested in all except the following two cases: (i) in the second set, mortar samples with Na_2SO_4 content up to 100% were also tested to allow understanding better the behaviour of this salt; (ii) in the case of the salt mixtures, samples with total salt content up to 5% were used because they were intended to represent salt mixtures found in real walls where 5% is already a very high salt content.

The deep narrow receptacles are small cylindrical plastic boxes with dimensions of about 5 cm in height by 3 cm in diameter. The shallow-wide receptacles are common glass Petri dishes of 10 cm in diameter.

A FITOCLIMA 500 EDTU® climatic chamber (by Aralab, Portugal) and an analytical weighing device with resolution of 10^{-4} g and accurate to 10^{-3} g were used. The technical specifications for the climatic chamber indicate a maximum RH fluctuation of ± 2% within the range of 10% to 98% RH. But in practice, over the period of the present experiments, maximum fluctuation of ±0.2% (±0.1% during around 70% of the time) was registered by the chamber sensor about the value of 95.0% RH.

3.5.3 - Results

3.5.3.1 - General

The results of the HMC experiments are presented in sections 3.5.4.2 to 3.5.4.9. In section 3.4.2.10 the error that would arise if the obtained HMC values were used to estimate the salt content in the samples is evaluated. When nothing is specified, the results refer to tests carried out on samples in deep narrow receptacles.

For most groups of samples (samples with different content of the same salt) regression lines are calculated that fit the obtained HMC values. The coefficient of determination (R^2) is a measure of how well the regression line represents the data and, therefore, allows evaluating whether a good linear correlation HMC / salt content exists.

The experiments were carried out with the climatic chamber set-pointed at 20°C and 80% or 95% RH. The HMC expected in equilibrium conditions at either RH is calculated by means of tables of thermodynamic data (Robinson and Stokes 2002, Robinson and Macaskill 1979, Roy *et al.* 1984). Such tables indicate the water activity a_w or the osmotic coefficient ϕ , by which a_w can be calculated through Eq. 3.1, for different molal concentrations of a given salt solution at a certain temperature. The molality *m* of the salt solution in equilibrium at 80.0% RH or 95.0% RH corresponds to water activity values of 0.80 or 0.95, respectively. The HMC expected for a sample with 100% salt content is calculated by merging equations 3.4 and 3.5 and then inserting the obtained value of *m*. The HMC of the base-materials is, hence, assumed to be zero.

Since thermodynamic data at 20°C, temperature at which the experiments were carried out, were only available for NaCl, data at 25°C were used: water activities for NaCl and KCl, osmotic coefficients for the other salts (Robinson and Stokes 2002, Robinson and Macaskill 1979, Roy *et al.* 1984).

For NaCl, water activities up to around 0.92 were available both for 20°C and 25°C by Olynyk and Gordon (1943). These values indicate that no HMC differences arise from that temperature difference, as seen in Fig. 3.11. However, this is not necessarily true for other soluble salts.



Fig. 3.11 - HMC of NaCl at 20°C or 25°C. Obtained from thermodynamic data by Olynyk and Gordon (1943).

The actual RH in the chamber is estimated using either:

- HMC of the pure salt samples;
- Regression equations of the contaminated brick or mortar samples. In this case, the HMC that corresponds to 100% salt content is considered.

These HMC values correspond to the concentration of the salt solution in equilibrium at the actual RH and temperature (Eq. 3.4 and 3.5). So, by linear interpolation on the tables of thermodynamic data, the water activity a_w of that solution can be determined. That water activity represents the actual RH in the chamber. Due to the lack of thermodynamic data at 20°C, data at 25°C is used

3.5.3.2 - Mass stabilization

In the experiments performed with deep narrow receptacles, mass stabilization of the samples could indeed be detected simply by visual assessment of the HMC versus time graphs. An example is presented in Fig. 3.12. The situation was different in the case of the shallow-wide receptacles as shown in the next section.



Fig. 3.12 - Evolution over time of the actual HMC of NaCl-loaded mortar samples in the deep narrow receptacles (second set of tests)

3.5.3.3 - Pure salts

Fig. 3.13 allows comparing the behaviour of the pure-salt samples in the two types of receptacles. This figure depicts the evolution over time of the HMC for the six salts whose RH_{eq}^{sat} is lower than 95% and for sodium sulfate. The experiments were performed at set-point conditions of 95% RH and 20°C. Fig. 3.14 allows evaluating agreement of the actual HMC values to the values expected in equilibrium conditions at 95.0% RH and 25°C (dotted lines).



Fig. 3.13 – Actual HMC of the pure salts in different receptacles: chlorides and nitrates (left); sodium sulfate and carbonates (right)



Fig. 3.14 – Comparison of the actual HMC of the pure salts in deep-narrow receptacles to the HMC expected at 95.0% RH and 25°C: chlorides and nitrates (left); sodium sulfate and carbonates (right).

These experiments on pure-salt samples were carried out in the second set of tests, as indicated in Table 3.1. Table 3.4 depicts the actual RH in the chamber during this second set, as estimated by means of the actual HMC of the salts. Two different moments were considered:

 the 29th day of testing which corresponds to the end of the tests on brick and mortar samples presented in sections 3.5.4.6 to 3.5.4.8; in this case, the HMC of the samples in shallow-wide receptacles was used because the samples in deep-narrow receptacles were still far from mass stabilization; the 64th day of testing which corresponds to the end of the tests on all pure salt samples; the HMC of the samples in deep-narrow receptacles was used in this case because it was in general more stable than that of the shallow-wide receptacles.

Table 3.4 - Actual RH during the second set: at the end of the tests on loaded brick or mortar (29 days) and on pure salt samples (64 days)

Moment at which the actual RH was estimated and samples used for that estimate	Actual RH in the climatic chamber (%) (estimated by means of the HMC of the pure salt samples)						
	NaCl	KCI	Na ₂ SO ₄	NaNO ₃	KNO ₃	Na ₂ CO ₃	K ₂ CO ₃
29 days (shallow-wide receptacles)	95.6	95.9	95.2	95.2	95.1	95.4	95.7
64 days (deep-narrow receptacles)	94.9	95.0	95.0	95.1	94.9	95.3	95.2

3.5.3.4 - Base-materials (mortar and brick)

The HMC of the samples composed only of brick (powder) or only of mortar (powder, granular or entire) is depicted in Fig. 3.15.



Fig. 3.15 - HMC of the base-materials

3.5.3.5 - Sodium chloride or sodium sulfate loaded materials: HMC at 80% RH

Fig. 3.16 depicts the HMC at set-point conditions of 80% RH and 20°C of brick and mortar samples with increasing contents of NaCl or Na_2SO_4 . In the figure, the dotted line corresponds to the HMC expected for NaCl in equilibrium conditions at 80.0% RH and 25°C. Table 3.5 presents, for NaCl, the equations of the regression lines that fit the experimental values and the estimates for the actual RH in the chamber. Since Na_2SO_4 attracted no moisture, no reference values are presented for this salt.



Fig. 3.16 - HMC of NaCl or Na₂SO₄ loaded mortar or brick samples at 80% RH

Table 3.5 - HMC at 80%RH and 20°C: actual RH and correlation equations for NaCl-loaded brick or mortar samples

Base- material	Line equation y - HMC (%) x - salt content (%)	Coefficient of determination (R ²)	Actual RH (%) at 25°C
mortar	y=3.4x+0.3	1.0000	80.5
brick	y=3.4x+0.1	1.0000	80.5

3.5.3.6 - Sodium chloride loaded materials

Fig. 3.17 depicts the HMC of NaCl-loaded brick or mortar samples at set-point conditions of 95% RH and 20°C in either set of tests. The dotted line represents the HMC expected at 95.0% RH and 25°C. Table 3.6 presents the linear regression equations that fit the experimental data and the corresponding estimates of the actual RH in the chamber.



Fig. 3.17- HMC of NaCl-loaded brick or mortar samples at 95% RH

Table 3.6 - HMC at 95%RH and 20°C: actual RH and correlation equations for NaCl-loaded brick or mortar samples

	Deee	Line equation	Coefficient of	Actual RH
Set	Base- material	y - HMC (%) x - salt content (%)	determination (R ²)	(%) at 25°C
1 st	mortar	y=12.6x+1.1	0.9998	95.4
1	brick	y=12.7x+0.3	0.9998	95.5
2 nd	mortar	y=13.1x+1.1	1.0000	95.6

3.5.3.7 Sodium sulfate loaded materials

Fig. 3.18 depicts the HMC of Na_2SO_4 -loaded brick or mortar samples obtained at set-point conditions of 95% RH and 20°C in either set of tests. Two different graphs are presented for reasons of legibility because a much larger range of salt contents was used for Na_2SO_4 than for the other salts. The dotted lines in this figure correspond to: (i) HMC values expected at

95.0% RH and 25°C; (ii) water (%W) that would be held by the samples if the thenardite initially present in these samples totally hydrated into mirabilite; (iii) MC corresponding to the maximum solubility of the metastable heptahydrate, obtained from solubility data at 20°C (Lurie 1975).



Fig. 3.18 - HMC of Na₂SO₄-loaded brick or mortar samples at 95% RH: higher (left) and lower (right) salt contents.

Table 3.7 presents the equations of the regression lines that fit the experimental data and the corresponding estimates of the actual RH in the chamber. Both the equations and the actual RH were calculated only for salt contents below 2% because, as seen in Fig. 3.18, the HMC obtained at higher salt contents is too erratic to fit a linear regression.

Set	Base-material	Line equation y - HMC (%) x - salt cont. (%)	Coefficient of determination (R ²)	Actual RH (%) at 25℃
1 st	mortar	y=4.0x+1.1	0.9992	94.3
	brick	y=4.9x+0.3	0.9994	95.3
2 nd	mortar	y=5.1x+1.1	0.9933	95.4

Table 3.7 - HMC at 95%RH and 20°C: actual RH and correlation equations for mortar or brick samples with Na₂SO₄-content below 2%

Aiming at a better understanding of the apparently random behaviour of the samples with Na_2SO_4 content above 2%, a more detailed analysis of the results obtained in the second set of experiments is presented bellow. That was possible because, after the unexpected behaviour of Na_2SO_4 in the first set, more frequent (weekly) weighting and visual observation of the samples were carried out in the second set.

Fig. 3.19 depicts the absorption dynamics of the Na_2SO_4 -loaded samples during the second-set experiments. Two samples with salt content above 50%, which were not presented in Fig. 3.18 because they had not achieved equilibrium at the end of the test, are also included here.



Fig. 3.19 – Evolution over time of the HMC of mortar samples of the second set of tests with Na_2SO_4 content up to 100%

Visual observation of the samples at the end of the second set experiments allowed the observation of the following alterations:

- all the samples with salt content above 10%, except the sample with 75% salt content, were covered with liquid;
- a solid crust of salt was present on the surface of the sample with 75% salt content;
- a large crystal immerse in the solution was seen for the 100% salt sample.

The samples with less than 10% of salt showed no significant changes.

3.5.3.8 - Mortar loaded with other soluble salts

Fig. 3.20 depicts the HMC of mortar samples with increasing content of NaCl, KCl, K_2SO_4 , NaNO₃, Na₂CO₃ or K_2CO_3 . These experiments were performed at set-point conditions of 95% RH and 20°C in the second set. The results of NaCl were previously presented in Fig. 3.17 and Table 3.6 but are repeated here to permit a comparison. Na₂SO₄ was not included due to its random behaviour. CaSO₄.2H₂O and KNO₃ are also not included because they had no significant absorption of moisture (HMC of around 1.2% was obtained in both cases). Table 3.8 presents the linear regression equations that fit the experimental data, as well as the corresponding estimates of the actual RH in the chamber.



Fig. 3.20 - HMC at 95% RH of mortar samples loaded with each of six distinct salts

Salt	Line equation y - HMC (%) x - salt content (%)	Coefficient of determination (R ²)	Actual RH (%) at 25°C
NaCl	y = 13.1x + 1.1	0.9999	95.6
KCI	y = 9.9x + 1.1	0.9997	95.8
NaNO ₃	y = 8.1x + 1.1	0.9998	95.8
KNO3	y = 5.3x + 1.1	0.9993	95.5
Na ₂ CO ₃	y = 7.7x + 1.1	0.9978	95.5
K ₂ CO ₃	y = 6.6x + 1.1	0.9966	95.3

Table 3.8 - HMC at 95%RH and 20°C: actual RH and correlation equations for mortar samples loaded with six distinct salts

3.5.3.9 - Salt mixtures

Fig. 3.21 presents the HMC values obtained for mortar samples with increasing content of each of the four mixtures of NaCl, NaNO₃ and Na₂SO₄ whose composition is depicted in Table 3.3. The experiments were carried out at set-point conditions of 95% RH and 20°C in the second set. Table 3.9 presents the equations of the corresponding regression lines.



Fig. 3.21 - HMC at 95% RH of mortar loaded with four different salt mixtures

Table 3.9 - HMC at 95%RH and 20°C: correlation equations for mortar samples loaded with four salt mixtures

	HMC of salt mixture loaded mortar (%)			
Mixture	Line equation	Coefficient of determination (R ²)		
Α	y = 8,0 x + 1,1	0,9998		
В	y = 9,6 x + 1,1	0,9992		
С	y = 9,8 x + 1,1	0,9995		
D	y = 8,8 x + 1,1	0,9993		

Fig. 3.22 allows comparing the actual HMC of the mixtures to the values calculated by weighing the HMC of the individual salts, either in the case of their actual HMC (Tables 3.7 and 3.8) or of the HMC expected at 95.0% RH and 25°C. Table 3.10 quantifies the agreement of the actual HMC of the mixtures to these weighted values.



Fig. 3.22 - Actual HMC for the salt mixtures and theoretical HMC calculated by weighing either the actual HMC or the HMC expected at 95.0% RH and 25°C of the individual salts

Table 3.10 - Agreement of the actual HMC of the	mixture with the value obtained by weighting
the HMC of the in	dividual salts

			Mixture A	Mixture B	Mixture C	Mixture D
Linear regression equations	Actual HMC of the mixture at 95% 20°C (%)		HMC _{act} ≈8.0 x	HMC _{act} ≈ 9.6 x	HMC _{act} ≈ 9.8 x	HMC _{act} ≈ 8.8 x
	HMC of the mixture (%) estimated by weighing the actual HMC of the salts at 95% and 20°C		HMC _{est} ≈8.8x	HMC _{est} ≈9.8x	HMC _{est} ≈10.6x	HMC _{est} ≈9.1x
	HMC of the mixture (%) estimated by weighing the HMC of the salts expected at 95.0% and 25°C		HMC _{est} ≈7.7x	HMC _{est} ≈8.6x	HMC _{est} ≈9.2x	HMC _{est} ≈8.1x
Significance of the difference between the actual and the estimated HMC of the mixture:		When the HMC of the mixture is estimated by means of the actual HMC of the individual salts	-9.1	-2.0	-7.5	-3.3
$100 \frac{HMC_{act} - HMC_{est}}{HMC_{est}} (\%)$		When the HMC of the mixture is estimated by means of the HMC expected for the individual salts	3.9	11.6	6.5	8.6

* For sodium sulfate, the linear equation corresponding to salt content of less than 2% (Table 3.7) was used

Table 3.11 allows evaluating the maximum dispersion of the HMC between the four mixtures. This is done by considering the maximum HMC difference, which is between mixture C and mixture A (Fig. 3.21), and then accessing its significance in relation to the average HMC of the four mixtures.

Table 3.11 - Significance of the maximum HMC variation among the salt mixtures

Mixture C : HMC ^{max} ≈ 9.8 x	HMC - HMC
Mixture A: HMC ^{min} ≈ 8.0 x	$\frac{100 \frac{1100 \text{ max}}{\text{max}} \text{ mme}_{\text{min}}}{\text{HMC}} = 19.9 \text{ (\%)}$
Average of the four mixtures: HMC ^{av} ≈ 9.1x	av

3.5.3.10 - Error in salt content estimates

Table 3.12 presents the error α (Eq. 3.9) that arises when the salt content in the samples is estimated, by means of the actual HMC values obtained at set-point conditions of 95% RH and 20°C, using as reference the HMC values expected at 95.0% and 25°C.

	Samples (Actual HMC at 95% RH and 20°C		HMC expected at	
Set		RH (%)	Equation y=HMC (%) x=salt content (%)	y(x=100%) concerns parameter E ⁺ in Eq. 3.9	95.0% and 25°C for samples with 100% salt content (concerns parameters $E^- \equiv E$ in Eq. 3.9)	$\alpha = 100 \frac{E^+ - E}{E} (\%)$ (from Eq. 3.9)
1	mortar + NaCl	80	y=3.4x+0.3	340.3	332.5	2.3
1	brick + NaCl	80	y=3.4x+0.1	340.1	332.5	2.3
1	mortar + NaCl	95	y=12.6x+1.1	1270.3	1153.4	10.1
1	brick + NaCl	95	y=12.7x+0.3	1261.1	1153.4	9.3
2	mortar + NaCl	95	y=13.1x+1.1	1310.3	1153.4	13.6
1	mortar + Na ₂ SO ₄ (<2%)	95	y=4.0x+1.1	400.3	463.6	-13.6
1	brick + Na ₂ SO ₄ (<2%)	95	y=4.9x+0.3	490.3	463.6	5.8
2	mortar + Na ₂ SO ₄ (<2%)	95	y=5.1x+1.1	511.1	463.6	10.3
2	mortar + KCl	95	y = 9.9x + 1.1	991.1	853.1	16.2
2	mortar + NaNO ₃	95	y = 8.1x + 1.1	811.1	692.0	17.2
2	mortar + KNO ₃	95	y = 5.3x + 1.1	531.1	459.1	15.7
2	mortar + Na ₂ CO ₃	95	y = 7.7x + 1.1	771.1	671.2	14.9
2	mortar + K ₂ CO ₃	95	y = 6.6x + 1.1	661.1	616.3	7.3

Fable 3.12 - Error ir	 salt content estimates
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3.5.4 - Discussion

3.5.4.1 - Influence of the base-materials and their state of cohesion

As expected, the HMC of the base-materials (mortar and brick) is extremely low (Fig. 3.15) when compared to the HMC of the soluble salts (Fig. 3.13). Accordingly, the base-material does not have a relevant influence on the HMC of the salt-loaded materials: materials with the same salt content result in similar HMC values, regardless of whether they are brick or mortar. This happens both for NaCl (Figs. 3.16 and 3.17) and for Na₂SO₄ (Figs. 3.16 and 3.18) whose HMC deviations are clearly not related to the nature of the base-material.

Further, no significant differences are found (Fig. 3.15) between the results obtained for the three cohesion states of the mortar samples (powder, granular and entire pieces). This corroborates the conclusions of other authors (Lubelli *et al.* 2004).

3.5.4.2 - Mass stabilization

The shape of the receptacle can significantly influence the mass stabilization of the samples. Fig 3.13 depicts the results obtained with the two types of receptacles and shows that:

- The absorption/dissolution process of the salts is slower when deep narrow receptacles are used. This is probably due to: (i) the low velocity of the air, thus, to the low moisture transfer rate between the sample and the adjacent air in these receptacles which are deeper than the Petri dishes; (ii) the smaller exposed area of the material/solution in these receptacles which are narrower than the Petri dishes. Accordingly, hygroscopic equilibrium takes longer to be reached in the deep-narrow receptacles than in the shallow-wide Petri dishes.
- In general, the HMC is more stable for the deep-narrow than for the shallow-wide receptacles. With the deep-narrow receptacles all the samples ultimately achieve a constant mass. But with the shallow-wide Petri dishes this does not occur for all the salts. Apparently, the deep-narrow receptacles, by slowing the moisture transfer rate, also "smooth" the effects of the environmental oscillation on the HMC.
- The HMC is more stable for some salts (KNO₃, NaNO₃ and even Na₂SO₄) than others (NaCl, KCl, Na₂CO₃ and K₂CO₃) particularly when shallow-wide receptacles are used. Perhaps a fast dissolution kinetics and high HMC is what determines the tendency of some salts for generating unstable HMC values. KNO₃, for example, which has the slower hygroscopic reaction and one of the lowest HMC, shows stable HMC values in both types of receptacles. By contrast, NaCl, which has a very fast hygroscopic reaction and the highest HMC, shows less stable HMC when shallow-wide receptacles are used. However, some situations are not totally clear. For example, the HMC and hygroscopic reaction of NaNO₃ and K₂CO₃ are quite similar. Yet, in the shallow-wide receptacles, the HMC of NaNO₃ is very stable, while the HMC of K₂CO₃ oscillates.

3.5.4.3 - Correlation HMC/salt content

Good linear correlations HMC / salt content are obtained, either in or among the two sets of experiments, for the salts whose RH_{eq}^{sat} is below the actual RH in the chamber. Figs. 3.17 and 3.20 and Tables 3.6 and 3.8 show good linear correlations HMC / salt content at 95% RH for NaCl, KCl, K₂SO₄, NaNO₃, Na₂CO₃ and K₂CO₃. The same occurs for NaCl at 80% RH, as shown in Fig. 3.16 and Table 3.5.

Sodium sulfate is a special case. Although the pure Na_2SO_4 samples achieve the equilibrium concentration (Fig. 3.14), the salt-loaded brick or mortar samples do not provide sound linear correlations. At 95% RH, a reasonable linear correlation is found only for Na_2SO_4 contents of 2% or less (Fig. 3.18). Further, at 80% RH, the Na_2SO_4 -loaded samples do not attract any moisture (Fig 3.16) despite the air RH being higher than that (77% RH) corresponding to the hydration of thenardite (Fig. 2.9).

With the four salt mixtures, good linear correlations are found (Fig. 3.21 and Table 3.9), nonetheless sodium sulfate is present in three of them.

Gypsum and potassium sulfate showed no significant absorption of moisture because their RH_{eq}^{sat} is of 97.6 % and 99.6 %, respectively (Table 3.2). Therefore, hygroscopic absorption is not expected to occur at 95% RH.

The good linear correlations HMC / salt content obtained for all salts (but Na_2SO_4) and salt mixtures suggest that the environmental conditions in the chamber were quite homogenous in space (Tables 3.5, 3.6 and 3.8). The fact that very similar correlations were obtained for NaCl among the two sets (Fig. 3.17 and Table 3.6) indicates that these conditions were also reasonably constant in time (from set to set).

3.5.4.4 - Accuracy

HMC values quite similar to those expected at thermodynamic equilibrium are obtained, either at 80% or 95% RH, for all the pure salt samples and nearly all salt-loaded brick or mortar samples (Fig. 3.14, 3.16 and 3.17). Na_2SO_4 is the only exception and, hence, will be discussed in more detail in the next section.

Because the HMC of NaCl is not relevantly affected by temperature (Fig. 3.11), the use of reference data at 25°C instead of 20°C is not in this case relevant. Therefore, the RH values estimated by means of NaCl (Tables 3.4, 3.5 and 3.6) are likely to correspond to the actual RH in the chamber:

- 80.5% or 95.4% to 95.5% at the end of the first set experiments
- 95.6% in the first month of the second set (at the end of the tests on salt-loaded mortar samples)
- 94.9% at the end of the second set (end of tests on pure salt samples)

These values indicate that:

- the RH in the chamber was a few decimals above 95.0% in the first set;
- the same happened probably during the first month of the second set; the other salts tested in this period also indicate a actual RH a fraction above 95.0% (Table 3.8); this suggests that the HMC of these other salts was not relevantly affected by, nevertheless unlikely, temperature deviations;
- in the second set, a RH drop of around 0.5% probably occurred between the 29th and the 64th days.

As seen in Table 3.12, the error that would arise if the salt content in the samples was estimated by means of the actual HMC values is low at the set-point RH of 80% (2.3% for NaCl) but higher at the set-point RH of 95% (from 7.3% to 17.2% for all salts but Na_2SO_4). In the case of Na_2SO_4 , the error is very uneven (from -13.6% to 10.3% at 95% RH) even though it was calculated considering only the samples with less than 2% salt content.

3.5.4.5 - Sodium sulfate

Sodium sulfate is an exception among all the tested salts, as pointed in the previous sections. Despite the pure salt samples achieved equilibrium (Fig. 3.14) many of the salt-loaded brick or mortar samples had an apparently erratic behaviour.

At the set-point RH of 80%, the NaCl-loaded samples achieved concentrations that correspond to an actual RH of 80.5% (Table 3.5). This demonstrates the soundness of the environmental conditions in the chamber. Yet, the Na₂SO₄-loaded samples did not attract any moisture (Fig. 3.16) despite the hydration RH of Na₂SO₄ (77%, as seen in Fig. 2.9) being clearly below that actual RH.

This behaviour of sodium sulfate is not in agreement with the results of Lubelli *et al.* (2004) who reported hydration of thenardite into mirabilite at 80% RH and 20°C. But it may be in agreement with theories arguing that the hydration mechanism of thenardite to mirabilite is in practice "through-solution hydration", that is, dissolution of thenardite followed by precipitation of mirabilite from a supersaturated solution (Charola and Weber 1992, MacMahon *et al.* 1992, Rodriguez-Navarro *et al.* 2000). Since dissolution of thenardite at 20°C happens only at 87% RH (Fig. 2.9), direct hydration of thenardite to mirabilite cannot probably occur at 80% RH.

At set-point conditions of 95% RH, reasonable agreement with the values expected in equilibrium at 95.0% RH was obtained for all the samples with salt content of 2% or less (Fig. 3.18 and Table 3.7). Yet, for salt contents above 2%, apparently random results arose:

- In the first set, the HMC of the samples with 5%, 10%, 30% or 50% salt content is far from agreeing with the values expected at 95.0% RH (Fig. 3.18). Particularly, the samples with 30% or 50% salt content crystallized as mirabilite and remained (undissolved) in that condition.
- In the second set, most of the samples follow approximately the 95.0% RH equilibrium line (Figs. 3.18 and 3.19). However, the samples with 5%, 75% or 100% salt content have lower HMC, though the 100% sample would possibly attain the 95.0% RH equilibrium line if the test had not ended before its mass was stabilized.

The HMC of the 75% sample in the second set and 5% samples in either set agrees approximately to the maximum solubility of the metastable heptahydrate (Figs. 3.18 and 3.19). In particular, the 75% sample attained a maximum HMC value at the 1.5 months, which afterwards started decreasing. This indicates that crystallization started at the maximum solubility of the heptahydrate and was still occurring at the end of the test. Therefore, it is probable that the phase observed at the surface of the 75% sample was the heptahydrate.

This random behaviour of Na₂SO₄ at the set-point of 95% RH arises probably from the actual RH in the chamber being, most of the time, practically coincident to the RH_{eq}^{sat} of mirabilite (Table 3.6, for instance). Indeed, the set-point RH of 95% was chosen based on the former assumption that the RH_{eq}^{sat} of mirabilite was of 93.6% at 20°C (Price and Brimblecombe 1994, for instance). But further theoretical work (Linnow *et al.* 2006) indicated a RH_{eq}^{sat} of 95.6% at 20° (Fig. 2.9). The present experimental results corroborate, therefore, this new value.

3.5.4.6 - Salt mixtures

All salt mixtures provided good linear correlation HMC / salt content, as seen in Fig. 3.21 and Table 3.9, despite sodium sulfate being present in three of them (Table 3.3).

The agreement with the HMC estimated by weighing the (actual or equilibrium) HMC of the individual salts is also very reasonable (Fig. 3.22 and Table 3.10). Indeed, the difference is always less than 12% of the estimated HMC. This suggests that, at 95% RH, the solutions become dilute enough for the salts in the mixtures not to have a very significant interaction. On the contrary, in strong aqueous solutions these interactions are very strong, hence, the behaviour of the mixture cannot be predicted on the basis of the behaviour of the individual salts (Price and Brimblecomb 1994).

In a previous study (Lubelli *et al.*, 2004) on a mixture of 50% Na₂SO₄, 25% NaCl and 25% NaNO₃ a reasonably good agreement was also found between the actual HMC of the mixture at 96% RH and 20°C and the HMC estimated by weighing the actual HMC of the three salts in similar environmental conditions. However, a significant difference was found between the actual HMC of the mixture and the HMC estimated by weighing the individual HMC values expected at equilibrium. Further, at 93% RH and 20°C, significant difference between the three HMC values was obtained. One of the possible causes that the authors indicate for these discrepancies is the actual RH being lower than expected. This seems probable, in accordance to what was argued in section 3.3, because the experiments were performed in closed containers where the RH was controlled by means of aqueous salt solutions.

The dispersion of the actual HMC values among the four mixtures is reasonably low: the maximum HMC difference is between mixtures A and C and represents around 20% of the average HMC of the four mixtures (Table 3.11). Though higher differences may exist for different salt mixtures, this suggests that estimates of the salt content sufficiently accurate for some practical purposes are perhaps possible by means of the HMC method, even if the salt mixture has some variation from sample to sample.

3.6 Conclusions

The overall conclusions of the work carried out to assess the possibilities and limitations of the two laboratory techniques, for RH control by salt solutions and salt content evaluation by the HMC method, are that:

- The use of aqueous solutions for RH control is not in general a safe method when salt-loaded materials are present.
- The HMC method can allow, although not in all situations, sufficiently accurate estimation of the salt content.

Use of aqueous solutions for control of the relative humidity

Aqueous solutions may not allow a stable control of the environmental conditions when salt-loaded materials are present. As demonstrated by the experimental research presented in section 3.3, both the RH and temperature may be disturbed during significant periods by secondary solutions that: (i) exist on wet materials; (ii) may form on dry materials when the RH in the containers is higher that the RH_{eq}^{sat} of the contaminant salt. Temperature disturbance is expected to be, in general, less significant over time because it arises predominantly from dissolution or crystallization processes. In contrast, RH disturbance may persist for longer periods following (slower) solution concentration changes.

Given the number and complexity of the possible influencing factors it is unfeasible to set general threshold values. Furthermore, the acceptable magnitude for the variation of the environmental parameters depends on the type of test. For instance, within HMC experiments, RH variability of only a few decimals may be already unacceptable. Indeed, as shown in section 3.4, the HMC of salt-loaded materials may suffer high magnitude deviations as a result of small RH variations. In contrast, in other tests, such as some crystallization tests, an RH range of several units may be acceptable. Therefore, case-by-case decisions should be made. Nonetheless, except for dry materials stored at an RH below the RH_{eq}^{sat} of the contaminant salt:

- Methods of controlling the RH by means of aqueous solutions should be avoided in the case of salt-loaded materials. The use of climatic chambers is preferable.
- If methods of controlling the RH by means of pure water or salt solutions are used on salt-loaded materials, the environmental conditions inside the testing containers should be monitored.

Evaluation of the salt content by means of the HMC method

The HMC method for salt content evaluation was systematized and discussed on the basis of theoretical and experimental work presented in sections 3.4 and 3.5, respectively. This research suggested that the method can be effective for most soluble salts when a RH above the $\rm RH_{eq}^{sat}$ of the salts is used.

Good linear correlations HMC / salt content and agreement with the values expected from equilibrium thermodynamic data were obtained for NaCl at 20°C and 80%RH or 95%RH, as well as, at 95%RH and 20°C, for KCl, K_2SO_4 , NaNO₃, Na₂CO₃, K_2CO_3 and the four mixtures of NaCl, Na₂SO₄ and NaNO₃.

Only for Na₂SO₄ a coherent linear correlation HMC / salt content could not be found:

- At 80% RH (20°C) that happened probably because the hydration of thenardite into mirabilite proceeds via dissolution of thenardite and further crystallization of mirabilite from a supersaturated solution. Therefore, since thenardite only dissolves at 87% RH, such process could not occur at 80% RH.
- At 95% RH (20°C), the random behaviour of sodium sulfate was most likely due to the actual RH in the chamber being practically coincident to the newly proposed RH_{eq}^{sat} of mirabilite which is of 95.6% (Linnow *et al.* 2006). In fact, the present experimental results could not be fully explained in a recent article (Gonçalves *et al.* 2006a) where the previous value of 93.6% RH was still considered. But they fit with and, hence, support this new value.

It is, therefore, recommendable to perform preliminary HMC experiments whenever possible, particularly when the behaviour of the involved salts at the chosen RH and temperature is poorly known. Nonetheless, in general, the HMC method is expected to allow either an absolute or at least a relative evaluation of the salt content.

Absolute determination of the salt content is possible:

- For salts of well-known thermodynamic properties, by comparing the actual HMC with the HMC expected at equilibrium (obtained from tables of thermodynamic data).
- For salts available/replicable at the laboratory, either their thermodynamic properties are known or unknown, by using an experimental correlation line obtained by means of control-samples.
- For some salt mixtures. At high RH such as 95%, very dilute solutions can form and, hence, the salts in the mixture do not have a significant interaction. This is suggested by the experimental results obtained here with NaCl, Na₂SO₄ and NaNO₃. For many practical applications, the HMC of mixtures with similar behaviour is sufficiently similar to the value calculated by weighing the (actual or equilibrium) HMC of the individual salts.

<u>Relative evaluation of the salt content</u> in several samples is theoretically possible, because the HMC is directly proportional to the salt content, for salts of unknown thermodynamic properties that are neither available nor are possible to replicate in the laboratory. That is the case of, for instance, samples collected from buildings which are normally loaded with complex salt mixtures. An estimate of the salt content sufficiently accurate for some practical purposes is still expected to be possible, even if the salt mixture is not totally homogeneous from sample to sample. Indeed, a maximum HMC variation of around 20% in relation to the average HMC of the four mixtures of NaCl, Na_2SO_4 and $NaNO_3$ was obtained, despite the significant variation of the relative contents of the three salts (Table 3.3). When performing HMC measurements, it must be considered that minor RH deviations may induce significant HMC variations. The higher the actual RH in the chamber, the higher HMC variations can arise from small RH deviations because the HMC of soluble salts increases exponentially with the RH (at 100% RH, the HMC tends to infinity). Therefore, the accuracy of the method depends largely on the capability of the climatic chamber to generate accurate and homogeneous environmental conditions in space and in time.

The following methods are proposed to reduce the uncertainty associated to salt content estimation by the HMC method:

a) Experimental correlation lines obtained by means of control-samples of the same salt, rather than theoretical values obtained from thermodynamic data in the literature, may be used as reference to calculate the salt content in the contaminated samples. So, permanent or erratic deviations from the set-point RH or temperature will not be a problem (unless the magnitude of those deviations is such that the RH_{eq}^{sat} of the salt is not reached). This method can be used in the case of salts available or replicable in the laboratory, regardless of whether their thermodynamic behaviour is known or unknown.

Only pure salt samples are required to obtain the correlation line because the HMC is directly proportional to the salt content. Control-samples of the base-material can be used as well but, in most cases, it is perfectly acceptable to assume the HMC of the base-material as zero. Indeed, the HMC of common building materials is insignificant when compared to the HMC of soluble salts, regardless of the state of cohesion of the sample.

- b) Control-samples of a known salt may be used to evaluate the agreement of the actual RH to the set-point RH, by comparing the actual HMC of these samples to the HMC expected at thermodynamic equilibrium. For this, the actual temperature in the chamber needs to be sufficiently accurate and constant, which is not normally a difficulty because temperature control in climatic chambers is relatively simple when compared to RH control. Furthermore, the sensitivity of soluble salts to temperature changes is in general much lower than to RH changes, particularly at high RH. For some salts, such as NaCl, the temperature is not at all relevant in the ordinary range of temperatures used in HMC experiments.
- c) When HMC values obtained by means of equilibrium thermodynamic data in the literature are to be used as a reference, either to estimate the salt content in the contaminated samples or to assess the actual RH, it is perhaps preferable to perform the experiments at a temperature for which such data is available, for instance 25°C rather than 20°C. Though, this is not necessary for salts like NaCl which are not sensitive to temperature.
- d) Several control-samples should always be used, spread throughout the chamber. The variability of their HMC allows evaluating the spatial homogeneity of the environmental conditions in the chamber. If that variability is significant, it may be necessary to place all the samples in a limited area of the chamber. If very different HMC values are still obtained, the climatic chamber is inappropriate for HMC determination.
- e) The control-samples should always be placed next to the loaded samples.

Non-despicable mass variations, which moreover increase linearly with the salt content in the sample, can arise also from small oscillations of the RH that ordinarily occur in climatic chambers. For that reason, the usual criterion of considering that equilibrium was achieved when the mass variation is lower than a certain percentage of the dry mass of the sample is

hardly applicable. In practice, hygroscopic equilibrium can be appropriately detected simply by observation of the HMC-versus-time graph.

Further, when deep narrow recipients are used instead of usual Petri dishes, hygroscopic equilibrium takes longer to be reached but the HMC fluctuation is smoothed. In the HMC measurements performed in this thesis, current Petri dishes were used in several cases (Chapter 4 and 5) and no major stabilization problems arouse. But, because the sensitivity of soluble salts to variations of the environmental conditions can be different, that will not necessarily happen for all climatic chambers, types of salt and salt contents.

Deep-narrow receptacles were used in this thesis to measure the HMC of samples collected from buildings (Chapter 6). One of the main reasons was the very large number of samples involved, which was of the order of several hundreds. The use of these small sized receptacles allowed testing many dozens of samples simultaneously and, hence, accomplishing the HMC measurements within an acceptable period.

Chapter 4 – Drying of salt-loaded materials

4.1 Introduction

Salt decay processes and features are very closely related to masonry drying. In fact, it is typically during drying that salts crystallize. Moreover, evidence exists that the so-called "salt-induced dampness" may arise from an alteration of the drying process that happens under the influence of soluble salts. But how does salt-loaded masonry dry?

The depth at which salts crystallize in porous materials depends on the balance between the liquid flux to the drying front and evaporation flux (Lewin 1982). Yet, some salts have a high tendency to form efflorescence while others, under similar environmental conditions, tend to deposit within the pores of building materials. For instance, in laboratory crystallization tests, which can be defined as drying experiments performed under particular conditions, sodium sulfate tends to form subflorescence, while sodium chloride typically produces abundant efflorescence (Goudie and Viles 1997, for instance). The reasons for this distinct behaviour require further explanation.

In buildings, some salts tend to crystallize higher on the walls, while others tend to be deposited closer to the ground. Arnold and Zehnder analysed the distribution of salt efflorescence on many monuments affected by rising damp (Arnold 1982, Arnold and Zehnder 1989) and observed that different types of salts appeared generally at distinct heights. The authors attributed this feature to the fractional crystallization of the salts from a mixed solution, a process that occurs sequentially according to the solubility of the individual salts. But a question arises whether in the (theoretical) case of similar walls contaminated with a single salt, the height of deposition would be similar for different salts.

In this approach, the materials are analysed on a large scale. However, masonry is a heterogeneous system. They are composed of distinct elements: plaster, bedding mortar and stone or brick. Furthermore, masonry materials may vary from building to building. Hence, it is also important to understand drying and deposition processes in more detail.

Indeed, the movement of liquids within heterogeneous system depends, to a large extent, on the suction properties of the individual materials in contact. For instance, pointing mortars should favour liquid transport towards the surface during drying in order to accomplishing their sacrificial role and, hence, protect the masonry elements. It is common knowledge that when, for example, water repellent or very dense pointing mortars are used, salt solutions are diverted towards the adjacent stone or brick elements which eventually decay. In old

plastered/rendered masonry that sacrificial role is traditionally attributed to the plaster/render. Lime plasters and renders were normally used because they favour salt crystallization on the surface or in their own mass, thereby preventing the underlying masonry from decaying. However, evidence of in-depth loss of cohesion of lime bedding mortars in old masonry, where only traditional lime plasters/renders exist, is often found in practice. Therefore, a question arises as to whether crystallization in old masonry can always and completely be prevented by using very absorbent plasters and renders that favour liquid migration to the surface.

Salt-loaded walls recurrently present moist surfaces. The height of capillary rise, namely, is typically higher when soluble salts are present (Massari and Massari 1993). However, as recently stated by an experienced building pathologist concerning the common case of rising damp, "... the examination of two closely affiliated phenomena, salt damage and rising damp, leaves us with far less in the way of definitive intervention strategies than we might wish, primarily because our understanding of the phenomena is limited" (Harris 2001). It is therefore crucial to understand the mechanisms of salt-induced dampness.

Hygroscopicity of soluble salts is a frequent cause of salt-damp in old buildings, as verified for instance by Henriques (2006). However, many of the salts currently present in walls have reasonably high relative equilibrium humidity (Table 3.2) and, hence, hygroscopicity can not probably explain all cases of salt-induced dampness.

Hygroscopic absorption occurs, as explained in section 3.2.3, when the air RH is higher than the RH_{eq} of the contaminant solution. From a thermodynamic point of view, a negative vapour pressure gradient exists in this case between the salt-loaded material and the surrounding air. Thus, water vapour diffuses from the environment to the surface and the material absorbs moisture. Differently, when the air RH is lower than the RH_{eq} of the salt solution, the vapour pressure gradient is positive. Therefore, water vapour diffuses in the opposite direction and the salt-loaded material dries.

By comparison to pure water, the driving vapour pressure gradient for vapour transport between a wet material and the environment is lower when soluble salts are present because non-volatile solutes depress the RH_{eq} of the solution. This is why porous materials dry more slowly if soluble salts are present. In consequence of that slower drying, the evaporation front tends to be located closer to the outer surface, as indicated by theoretical considerations of Hall and Hoff (2002). This suggests that salt-induced dampness may arise, in general, from changes in the drying process of porous building materials.

Aiming at a better understanding of these processes, drying experiments were carried out on specimens composed of stone, bedding mortar and plaster (Gonçalves *et al.* 2006c). The distribution of moisture during drying was monitored by means of the two-dimensional (2D) magnetic resonance imaging (MRI) technique presented in Annexe I. The technique provides sequences of images which map the concentration of liquid water in the specimens at given time intervals. MRI techniques are powerful tools whose use on building materials is still very restricted. In fact, to the best of this author's knowledge, this was the first time that a 2D MRI technique was used to investigate soluble salts in building materials. This initial study was restricted to pure water and NaCl contamination, as well as to isothermal drying conditions.

The main objectives of this research were then:

- To understand the influence of soluble salts on the drying of porous building materials;
- To explain the general mechanism of salt-induced dampness;
- To discuss the distribution of salt during drying in walls contaminated with a single salt, both at the (larger) scale of the wall and at the (smaller) scale of the masonry elements.

The work carried out is presented in this chapter. Fundamentals of moisture transport and drying of porous materials are introduced in section 4.2. The materials and methods used in the experiments are described in section 4.3. The experimental results are presented and discussed in sections 4.4 and 4.5, respectively.

4.2 Drying of porous materials

4.2.1 - Porosity and hygroscopicity

Building materials such as stone, brick, mortars or concrete contain small voids that are called "pores". For this reason, these materials are currently called "porous building materials". A percentage, usually significant, of these pores is interconnected, hence forming an internal network through which water migrates either as liquid or vapour. The open porosity of a material, often simply named "porosity", expresses the ratio of the total volume of interconnected pores to the total volume of the material. These interconnected pores are ordinarily classified into three main types, according to their size and resulting influence on the moisture transport properties of the material:

- The micropores are the smallest pores. They have no influence on moisture transport because the water they contain is not free water. Due to their very small size, the capillary forces are so high that, in current conditions, the water cannot move.
- The mesopores are pores of intermediate size. They are also called "capillary pores" or simply "capillaries". It is through them that liquid water migrates under the influence of capillary forces. Vapour transport also occurs in these pores.
- The macropores are the largest pores. They are very relevant for vapour transport. In contrast, liquid capillary transport is not normally significant in the macropores because the capillary forces are weak.

The size attributed to each type of pore is roughly similar among different authors. Hilbert *et al.* (1992), for instance, adopted the following classification during research on restoration plasters:

- micropores: radius less than 0.1µm;
- capillary pores: radius in the range of 0.1μm to 30μm (a sub-class of "highly capillary" pores with radius from 0.1μm to 15μm was also considered);
- macropores: radius larger than 30µm.

Most porous building materials such as stone, concrete, mortars or brick are hygroscopic materials. "Hygroscopic" is a general term that refers to substances that are able to attract water vapour from the environment. The hygroscopic behaviour of soluble salts was discussed in detail in section 3.2.3. In the case of building materials, hygroscopicity concerns adsorption of water from the air. Adsorption is the process by which water molecules bond with the pore surfaces and thereby forms thin films, composed of one or several layers of molecules. Adsorption arises from weak intermolecular (Van der Waals) forces that act between the material surface and water. The thickness of the adsorbed water films increases with the air RH. This is why, at a given temperature, the equilibrium HMC of porous building materials depends on the RH.

4.2.2 - Liquid water transport

In porous building materials, liquid water migrates mostly by capillarity. The mechanism of capillarity takes place when the attractive forces between the liquid and the solid material are stronger than the cohesive forces in the liquid.

The driving pressure for capillary suction, also called capillary pressure or suction stress (Künzel 1995), is given by Jurin's law which concerns an idealized cylindrical capillary (Eq. 4.1 and Fig. 4.1). In this equation, p_c (Pa) is the capillary pressure, σ (N/m) the surface tension of the liquid, r (m) the radius of the capillary and φ the contact angle.



Fig. 4.1 – Capillary pressure in a cylindrical capillary

The contact angle is the angle at which the liquid surface meets the solid surface. It results from the equilibrium of the tensions at the solid-vapour, solid-liquid and liquid-vapour interfaces. The contact angle between water and a hydrophobic surface is greater than 90°. The contact angle between water and a hydrophilic surface, for instance the pore walls of ordinary building materials, is less than 90°, which corresponds to a concave meniscus and to a negative capillary pressure, hence, to capillary movement of water into the material.

In a cylindrical capillary where the contact angle φ is less than 90° (Fig. 4.1), water rises up to where the capillary pressure is equilibrated by the weight of the water column. Where *p*

 (N/m^2) is the pressure generated by the water column, ρ (kg/m³) the density of water, g (m/s²) the gravitational constant and H_0 (m) the height of capillary rise in equilibrium conditions:

$$\rho = -\rho \cdot g \cdot H_0 \tag{Eq. 4.2}$$

Combining Eq. 4.1 and Eq. 4.2, the height of capillary H_0 rise is obtained as a function of the pore radius (Eq. 4.3):

$$H_0 = \frac{2 \cdot \sigma}{r \cdot \rho \cdot g} \cdot \cos \varphi \tag{Eq. 4.3}$$

This means that both the suction stress (Eq. 4.1) and the height of capillary rise (Eq. 4.3) vary in inverse proportion to the capillary radius.

The displacement of the water in a capillary is given, as a function of time, by Eq. 4.4 which derives from Poiseuille's experimental law (Washburn 1921):

$$d(t) = \sqrt{\frac{\sigma \cdot r \cdot \cos \varphi}{2\eta} \cdot t}$$
(Eq. 4.4)

In this equation d (m) is the distance which the meniscus travels in the capillary during time t (s), η (kg.m⁻¹.s⁻¹) the dynamic viscosity of water, σ (N/m) the surface tension of water, r (m) the capillary radius and φ the contact angle. It shows that water travels more quickly through larger capillary pores which, hence, are filled first during soaking.

When a porous material is in free contact with water it soaks liquid up to certain critical moisture content, the so-called "capillary saturation moisture content". Total saturation, which corresponds to total filling of the open porosity, cannot usually be reached under normal suction conditions within a reasonable period of time because a certain volume of air is generally trapped in the pore network, as well as because capillary suction by the macropores may be insignificant. In this thesis, the broad-sense term "saturation" refers to capillary saturation.

The capillary movement of water in porous materials cannot be predicted by means of the previous equations. In such materials, the capillary network is so complex that an individual analysis of the capillaries is not possible. A macroscopic approach is necessary. Indeed, liquid transport in porous materials is well described by means of an extended form of Darcy's equation. Unlike the original law which concerns saturated flow -in porous media, this extended law (Eq. 4.5) is applicable to unsaturated flow, the common case in buildings.

$$\vec{u} = -K_1 \cdot \vec{\nabla} p_c \tag{Eq. 4.5}$$

In this equation, \vec{u} (kg.m⁻².s⁻¹) is the density of the liquid flux, p_c (Pa) the capillary suction and K_1 (kg.m⁻¹.s⁻¹.Pa⁻¹) the unsaturated permeability coefficient or simply liquid conductivity which is a function of the moisture content.

The capillary transport properties of porous building materials are, consequently, also evaluated at the material scale. Appropriate experimental parameters are used to express the tendency of the material to, under specific conditions, absorb a certain amount of water or transport that water up to a certain height.

In a semi-infinite volume of a porous material in free absorption (Fig. 4.2), both the total amount of absorbed water and the height of capillary rise are direct functions of the square root of time. This is expressed by Eq. 4.6 and Eq. 4.7, respectively, which derive from Washburn's equation (Eq. 4.4).

$$W(t) = A \cdot \sqrt{t} \tag{Eq. 4.6}$$

$$H(t) = B \cdot \sqrt{t} \tag{Eq. 4.7}$$

In these equations, W (kg/m²) is the amount of absorbed water and H (m) the height of capillary rise after the period of immersion t (s). A (kg.m⁻².s^{-1/2}) and B (m/s^{1/2}) are the so-called "water absorption coefficient" (WAC) and "capillary penetration coefficient", respectively. These coefficients are complementary. A material can quickly absorb a high amount of moisture while presenting a low height of capillary rise and vice-versa. In fact, as seen before, smaller pores absorb the water more slowly but present higher heights of capillary rise at equilibrium (Eq. 4.3). In current language, the term "highly absorbent material" corresponds normally to a high WAC.

Coefficients *A* and *B* depend on the initial moisture content in the material, even though in practice they are typically measured on oven-dried materials. Indeed, in equilibrium conditions at given moisture content, all the pores below a certain diameter are filled with water. Capillary pressure varies in inverse proportion to the square root of the capillary radius (Eq. 4.1) and porous building materials contain pores of various sizes. Therefore, the sorptivity of a porous material, that is, its tendency to absorb and transmit liquids by capillarity (Hal and Hoff 2002), varies with its moisture content. This is compatible with the trivial observation that building materials absorb less water when they contain a higher amount of moisture.

In practice, the WAC is used much more than the penetration coefficient. One of the reasons is that direct measurement of coefficient B is often difficult and not very accurate. The boundary between wet and dry areas may not be easy to distinguish, particularly in micro- or macro-porous materials, or may be very irregular in composite or non-homogeneous materials. Sometimes, the penetration coefficient is estimated by dividing the WAC by the capillary saturation moisture content (Künzel 1995). This estimate is based on the assumption that the moisture content in the wet volume of the material is homogenous and identical to the capillary saturation moisture content. But this is not normally true and, so, the estimated heights of capillary rise are smaller than the actual ones.

In capillary absorption tests, specimens of regular dimensions and constant section, usually $4 \times 4 \times 16$ cm prisms, are allowed to freely absorb water through their bottom face (Fig. 4.2). The WAC is measured by weighing the specimens periodically, normally at increasing time intervals. Fig. 4.3 depicts two types of curves typically obtained, where the amount of absorbed water is plotted against the square root of time. The WAC corresponds to the inclination of the initial straight part of either curve.


Fig. 4.2 - Porous material in free absorption



Fig. 4.3 – Typical water absorption curves for porous building materials

Curve (a) in Fig. 4.3 is typical of materials whose pores have a limited range of dimensions, are well interconnected and homogenously distributed. A sharp transition between the wet and dry zones, that is, a well-defined dry-wet boundary exists in such materials during capillary absorption. The absorption curve is typically composed of two straight-line segments: (i) the first has a higher slope and corresponds to the filling of most of the interconnected porosity; (ii) the second has a lower slope and represents the further filling of the remaining pores by diffusion in water of the air trapped in the pore network. The inflection point corresponds to the moment when the capillary fringe reaches the top surface of the specimen. The ratio between the slopes of the two lines is generally within the range of 10 to 10^3 (Mertz 1991).

Curve (b) is typical of materials that contain several families of interconnected pores with clearly distinct dimensions. The different slopes reveal the existence of pores with different absorption kinetics (Mertz 1991, for instance). This behaviour corresponds, in the case of homogenous materials, to a diffuse dry-wet boundary during capillary absorption. In that case, the first inflexion point corresponds to the moment when the edge of that diffuse boundary reaches the top face of the specimen. The other inflexion points represent the moment when a given family of pores becomes saturated. Curve (b) can also occur for heterogeneous materials, for instance, composed by several horizontal layers of different (homogenous) materials. In this case, the inflexion points correspond to the moments when the capillary fringe reaches each of the interfaces between different materials.

Absorption curves with shapes different from those in Fig. 4.3 are sometimes obtained but they normally correspond to intermediate or even similar absorption behaviours. For example, a round-shaped absorption curve corresponds to a type (b) curve composed of many short segments.

The considerations above refer to ordinary hydrophilic materials. For materials that contain water repellent additives, such as most industrial plasters and renders, capillary absorption curves are interpreted in a similar way but in terms of capillary suction rather than pore dimension.

Experimental absorption curves usually intersect the ordinate axis somewhat above the origin, as shown in Fig. 4.3. Indeed, for a more accurate description of the absorption behaviour up to the first inflexion point, Eq. 4.6 could be written as Eq. 4.8 (Eq. 4.7 would undergo a similar change).

$$W(t) = A \cdot \sqrt{t} + w_0$$
 (Eq. 4.8)

Positive intercept w_0 in Eq. 4.8 arises because hydrostatic pressure adds to capillary pressure in the first mm of the specimen (below water level). A second possible cause is some water being absorbed through the lateral faces in the case of non-sealed specimens (Wilson *et al.* 1999, for instance)

In porous hygroscopic materials, liquid transport may occur even when there is no external contact with liquid water, particularly, when the material is exposed to an RH gradient. Indeed, the thickness of the water molecule layer adsorbed by the pore walls increases with the RH, hence, it may vary across a material exposed to an RH gradient. As a consequence, a (liquid) diffusion process takes place by which the water migrates from the thicker to the thinner adsorbed water layers. This process is called "surface diffusion" and is further illustrated in Fig. 4.4 d. Capillary transport of moisture may also arise in consequence of capillary condensation. Capillary condensation happens when the thickness of the adsorbed water layer increases to the point where the smaller capillaries are filled with liquid. Because of the RH gradient, the moisture content due to capillary condensation varies across the material and, hence, capillary transport may take place.

The pathways of water within systems composed of different materials are not easy to predict, even if hygroscopicity is not considered. This was well explained 70 years ago by Schaffer (1932 pp 51-53):

When different materials are placed in contact, a uniformly fine-pored material will draw water from a uniformly large-pored material, because the capillary suction is greater in small pores than in large ones. But building materials contain pores of various sizes, and in consequence the suction force developed will vary with the moisture content. In fact, the relative pore structures may even be such that the direction of the movement of water between two materials alters with change in the degree of saturation. (...) Since the capillary suction of a material varies with its moisture content, and the rate of change of moisture content under wetting or drying conditions varies with different materials, the problem of predicting the direction of transfer of water between any two materials is extremely complex.

Moreover, the interface between different materials in contact often introduces a resistance to moisture transport that is not easy to predict or quantify (Freitas 1992). The interface between two mortar layers, when one was directly applied to the other, represents perhaps the most favourable continuity condition as regards masonry materials. However, even this type of interface may not correspond to a situation of hydraulic continuity. This was, for instance, observed in previous work (Gonçalves 1997) where a significant hydric resistance was detected between two mortar layers due to the incomplete interpenetration of the two porous structures.

4.2.3 - Water vapour transport

Water vapour transport in porous building materials can be described as a diffusion process, hence, caused by a water vapour concentration gradient. In steady state isobaric and isothermal conditions, Fick's first law is applicable:

$$\vec{j} = -D_v \cdot \vec{\nabla}c \tag{Eq. 4.9}$$

In this equation, \vec{j} (kg.m⁻².s⁻¹) is the diffusion flux, *c* (kg/m³) the concentration of water vapour and D_v (m²/s) the diffusion coefficient or diffusivity.

Assuming that air behaves as an ideal gas, which is expressed by Eq. 4.10, the diffusion flux may be expressed as a function of water vapour pressure (Eq. 4.11).

$$p_{v} = \frac{c_{v} \cdot R \cdot T}{M_{w}}$$
 (Eq. 4.10)

Here, p_v (Pa) is the water vapour pressure, T (K) the temperature, R (J.K⁻¹.mol⁻¹) the ideal gas constant, c_v (kg/m³) the vapour concentration and M_w (kg/mol) the molar mass of water.

$$\vec{j} = -\Pi \cdot \vec{\nabla} p_v \tag{Eq. 4.11}$$

In this equation, Π (kg.m⁻¹.s⁻¹.Pa⁻¹) is the water vapour permeability of the material. Π expresses the quantity of water vapour crossing, in equilibrium isothermal conditions, a material of unitary thickness (m), per unit time (s), unit surface (m²) and unit difference of vapour pressure (Pa). Π depends on the temperature and water content of the material. Eq. 4.12 relates Π to the water vapour resistance μ .

$$\Pi = \frac{\delta}{\mu}$$
(Eq. 4.12)

Here, δ (kg.m⁻¹.s⁻¹.Pa⁻¹) is the coefficient of water vapour diffusion in air. Water vapour resistance μ (dimensionless) is a parameter that does not depend either on temperature or even, at low moisture contents, on the moisture content. Indeed, at higher moisture contents, the presence of liquid water in the pores may accelerate vapour transport because the liquid shortens the effective path length for vapour diffusion. This process is named "liquid-assisted vapour transfer" (Rose 1963a, 1963b) and is further depicted in Fig. 4.4 d.

Building materials are generally characterized, as regards vapour transport, by means of the well-known cup test methods. In these tests, an isothermal steady state of unidirectional vapour transport is induced by placing a specimen with certain thickness above a cup where a selected salt solution is used to generate a constant RH. The assemblage is placed in a room or chamber at constant RH and temperature. When steady state conditions are reached, the weight loss of the assemblage, which corresponds to the vapour flux through the specimen, is constant over time.

"Dry cup" or "wet cup" methods can be used where the RH in the cup is, respectively, lower or higher than the RH at the exterior. These methods are based on Eq. 4.13, which derives from Eq. 4.11 and expresses one-directional diffusion flux j (kg.m⁻².s⁻¹) that crosses a porous material when a given vapour pressure difference $p_{vi}-p_{ve}$ (Pa) is maintained and assumed to vary linearly across a specimen with thickness e (m).

$$j = \Pi \cdot \frac{(p_{vi} - p_{ve})}{e}$$
 (Eq. 4.13)

Besides Π (Eq. 4.13) and μ (Eq. 4.12) other quantities often used in practice are water vapour permeance Π/e (kg.m⁻².s⁻¹.Pa⁻¹) and diffusion-equivalent air layer thickness *s* (m). The latter is given by Eq. 4.14 and corresponds to the thickness of an air layer with the same permeance as the tested specimen.

$$s = \Pi^{air} \cdot \frac{p_{vi} - p_{ve}}{j} - s'$$
 (Eq. 4.14)

In this equation, Π^{air} (1,95 x 10⁻¹⁰ kg.m⁻¹.s⁻¹.Pa⁻¹) is the diffusion coefficient of vapour in air at atmospheric pressure and *s*'(m) is the thickness of the air layer in the cup (between the salt solution and the specimen).

Eq. 4.13 is used to characterize building materials in general. However, this equation is not strictly valid for hygroscopic materials because: (i) adsorbed water films can reduce the effective capillary radius; (ii) liquid transport by "surface diffusion" (Fig. 4.4 d) may arise simultaneously due to the vapour pressure gradient; (iii) capillary condensation may occur; (iv) liquid diffusive transport of that condensed water may take place due to the moisture concentration gradient; (v) vapour transport can be accelerated across water-filled pores (Fig. 4.4 d).

4.2.4 - Drying of porous materials

Wetting of porous materials from complete dryness is currently described, at the pore scale, by the six-phase process depicted in Fig. 4.4 (Rose 1963b). Each phase corresponds to different moisture content. Drying corresponds to the inverse process, nonetheless possible hysteretic effects that may arise in hygroscopic materials from a reduction of the effective capillary radius by adsorbed water layers.

At a macroscopic scale, drying of porous materials can be described as the three-phase process depicted in Fig. 4.5. The three drying stages, among which stages I and II are usually considered the most relevant (Sherwood 1929, Scherer 1990), occur as follows:

Stage I - Initially, the material is saturated and, hence, a continuous liquid phase exists in the pore network. Liquid water is transported towards the outer surface mostly by capillarity (Fig. 4.4e or 4.5f), though surface diffusion may also take place at the lower moisture contents (Fig. 4.4 d). A sharp capillary pressure gradient in the pores at the surface is the driving potential for capillary transport. Evaporation occurs at this surface and the moisture content in the material decreases at a constant rate. It is currently accepted that Stage I drying rate depends only on external factors such as the air RH, temperature and air velocity. Homogeneous moisture distribution is expected to exist throughout the material during this stage.

- Stage II When the unsaturated capillary flow is no longer able to supply water at a rate high enough to compensate evaporation, the drying front recedes into the material and drying Stage II starts. Liquid capillary transfer or vapour transport prevail beneath or ahead of the drying front, respectively. Receding of the drying front corresponds to the breaking of the liquid continuity in the pores close to the surface, which was argued to happen somewhere between phases (c) and (e) in Fig. 4.4 (Rose 1963b). The drying front goes on receding as evaporation proceeds and, hence, the moisture content in the wet zone decreases further. The drying rate of the material decreases over time with the lower availability of liquid at the drying front and with increase of the path length for vapour diffusion across the material. The drying front is sharp for materials with pores within a limited size range. When pores of significantly different size exist, the drying front is also expected to exist beneath the drying front during Stage II.
- Stage III The moisture content in the wet zone of the material decreases progressively during Stage II. Hence, at a certain moment, liquid continuity is broken in the capillaries beneath the drying front and drying Stage III begins. From this moment on, vapour diffusion controls the transport of moisture all over the material. Therefore, the drying rate becomes much lower. In this stage, although the material may be technically dry, clusters of liquid persist for some time in the smaller pores (Fig. 4.4 c). The drying rate is expected to decrease slowly, while the material approaches hygroscopic equilibrium (Figs. 4.4c to 4.4a).



Fig. 4.4 – Moisture transport at the pore scale during wetting of a porous material. After Rose (1963b).

Fig. 4.5 - Drying of a porous material at a macroscopic scale

The drying behaviour of porous building materials is normally accessed by means of the so-called "drying tests" or "evaporation tests" (RILEM 1980, for instance). In such tests, specimens of regular dimensions, typically prisms or cubes, are led to standard saturation conditions (capillary or even total vacuum saturation) with pure water. Afterwards, the lateral and bottom faces of the specimen are sealed in order to induce one-dimensional migration of moisture towards the top surface during drying. The specimens are allowed to dry under specific environmental conditions. By periodical weighing, a drying curve is obtained where either the actual moisture content in the material (Fig. 4.6) or the total amount of evaporated water is plotted against time. The drying rate of the material corresponds to the slope of the drying curve.



Fig. 4.6 – Typical drying curve of a porous building material

The transition between Stage II and Stage III is often indistinct in drying curves. In contrast, the transition between Stage I and Stage II is normally clear (Fig.4.6). It corresponds to the inflexion point between the straight (constant drying rate) and the round-shaped (decreasing drying rate) segments. The moisture content that corresponds to this inflexion point is named the "critical moisture content". That critical moisture content is typically higher for materials with less homogeneous pore size distributions (Snethlage and Wendler 1997). That happens because, when liquid continuity is interrupted in the larger pores, the smaller pores still contain a significant amount of moisture.

It is widely accepted that the drying rate of a material during Stage I depends on the external conditions. Yet, the inclination of the straight part of the drying curve may vary among different porous materials under similar evaporation conditions. This happens because different porosities may lead to different effective surfaces of evaporation in specimens of similar dimensions, hence, to different drying rates. Further, it has recently been suggested that also the pore size could influence the effective surface of evaporation. Rousset-Tournier (2001) observed that the drying rate of building stones during Stage I was systematically higher than the drying rate of a flat surface of water under identical environmental conditions with no air velocity. This suggested that the curvature of the menisci at the drying front could increase the effective surface of evaporation.

Drying curves are normally obtained by gravimetric methods, as explained above. But it is also possible to use other methods of monitoring the moisture content during drying. In this chapter, drying curves obtained by means of MRI measurements will be presented.

4.3 Materials and methods

4.3.1 - Materials

Aiming at a better understanding of the drying process of salt-loaded masonry, MRI-monitored drying experiments were performed on specimens composed of three materials representing the three main elements of old stone masonry: stone, bedding mortar and plaster (Fig. 4.7).



Fig. 4.7 - Pictures of one specimen: (a) top view; (b) lateral view

Two different specimens were used. Each specimen included a different type of stone: stone D (specimen DL1) and stone M (specimen ML1). Both are very common stones in Portugal so they are expected to be representative of old ordinary masonry in this country, normally built from locally available materials (Segurado 1908). In both specimens, plaster and bedding mortar were made from the same lime mortar L. Table 4.1 and Fig. 4.8 show the composition and properties of the three materials in the specimens.

Prior to making the specimens, the cut surfaces of the stone elements were sandpapered in order to improve mortar/stone adherence. Next, bases composed of mortar and stone were made by applying lime mortar L on either stone D or M. The bases were kept in a conditioned room at 20°C and 65% RH for two weeks. Afterwards, the plaster was made by applying lime mortar L on these bases. In both cases, the mortar was applied: (a) after prewetting the surfaces; (b) with the help of acrylic plaques placed laterally during application and removed around 15 minutes later. The specimens were maintained in the same conditioned room during the following five months before testing. At the two months of age, they were laterally sealed with epoxy resin.

Material	Volumetric composition / nature	Porosity accessible to water (% V)		Capillarity
		capillary saturation	total (vacuum) saturation	(kg.m ⁻² .h ^{-1/2})
Lime mortar L	1:1.5:1.5 (dry hydrated lime: sand from the Tagus river: yellow pit sand from Corroios)	17.6	28.7	11.9
Stone D	Calcitic calcareous stone with fossil fragments (medium porosity)	11.2	15.4	3.9
Stone M	Calcitic calcareous stone (high porosity)	26.6	32.4	11.6

Table 4.1 – Composition, porosity and capillary coefficient of the materials



Fig. 4.8 – Pore size distribution (MIP)

The porosity accessible to water was measured on specimens of irregular shape (around 2 cm x 2 cm x 2 cm). The measurements were carried out both at total saturation and at capillary saturation. In the first case, the tests were performed according to RILEM procedure I.1 (RILEM 1980) by means of vacuum saturation. In the second case, a similar procedure was followed but saturation was obtained by simply immersing the specimens in pure water during an overnight period.

Unidirectional capillary absorption tests were carried out on 50 mm x 50 mm x 20 mm specimens whose lateral faces were sealed with an epoxy resin. First, the specimens were oven-dried at 60° C. Afterwards, they were allowed to freely absorb pure water through their 50 mm x 50 mm bottom face. That bottom face corresponds, in the case of the mortar specimens, to the contact surface with a brick substrate onto which the mortar had been applied and hardened. The specimens were weighed periodically, the excess water being removed by means of quick touches with a soft paper. During the absorption periods the specimens were kept inside a closed acrylic box and the free water level was maintained 5 mm above the bottom face of the specimen. Three specimens of each material were used.

The pore size distribution was measured by mercury intrusion porosimetry (MIP) according to LNEC's procedure LERO PE15, which is based on ASTM D 4404-84 (ASTM 2004). In MIP techniques the pore sizes are estimated by means of Jurin's law (Eq. 4.1), hence, the pores are assumed to have a cylindrical shape.

4.3.2 - Drying experiments

Either specimen was subjected to two MRI-monitored drying experiments, the first with pure water and the second with a 3 molal NaCl solution. The specimens were led to capillary saturation by total immersion for around two hours. This period was enough to reach constant mass at a precision of 0.01g. Afterwards, the bottom face of the specimens was sealed with Parafilm M® plastic film (the lateral faces were already sealed with epoxy resin).

The 2D MRI technique used is presented in section I.3 (Annex I). MRI images that map the concentration of liquid water in the specimens were collected every 30 min during drying. Drying was induced by air flow at 0% RH (temperature $\approx 18^{\circ}$ C), which was turned on after the initial image was obtained. For pure water, drying was monitored for 20 hours. For the NaCl solution, drying was monitored for 60 hours (specimen DL1) or 35 hours (specimen ML1).

Evaporation experiments were performed also with free liquid surfaces in order to obtain complementary background information. The evaporation rates of free surfaces of pure water and NaCl saturated solution were measured under environmental conditions of either 20°C and 50% RH or 20°C and 15% RH. Identical Petri dishes of 77 mm internal diameter and 18 mm depth were used. Two dishes of each liquid were tested in both cases. Drying took place in a climatic chamber with a low air velocity, the FITOCLIMA 500 EDTU® chamber by Aralab (Portugal) used in the HMC experiments presented in Chapter 3. The evaporation area of the liquids was possibly smaller than the internal area of the Petri dish due to hindering of air movement by the lateral walls of the dishes. However, comparative analysis is possible because an identical height of liquid (13 mm) was used in all cases.

4.3.3 - Salt distribution

The final distribution of salt in the specimens was determined by means of the HMC method described in Chapter 3 using common Petri-dishes.

After oven drying at 40°C and further removal of loose efflorescence, the specimens were cut longitudinally into two halves. Afterwards, by progressively disaggregating them on sandpaper, one half was divided in the vertical direction and the other in the horizontal direction. Hence, salt distribution could be evaluated along two perpendicular directions.

The samples thereby obtained were oven dried at 40°C. Then, they were kept inside a climatic chamber set-pointed at 20°C and 95% RH until hygroscopic equilibrium was achieved, that is, until their mass remained constant over time. The same FITOCLIMA 500 EDTU® climatic chamber was used.

Ten control samples of pure NaCl, as well as two non-contaminated samples of each base material (mortar and stones) were simultaneously tested. Since the HMC varies linearly with the salt content, the salt content in the samples could be calculated in relation to an actual

calibration line defined by the HMC of the base materials (0% of salt) and of the control samples (100% of salt). Average values of the HMC of each type of control-sample were used for this purpose.

As argued in Chapter 3, when discussing the scope and precision of the HMC method, minor variations of the environmental conditions (in particular of RH) may cause the concentration of salt solutions to vary. In the present HMC measurements, this uncertainty was evaluated by means of the ten NaCl control samples which were distributed throughout the chamber. Average, minimum and maximum HMC of respectively 732.2%, 688.3% and 756.4% were obtained among these samples. This values correspond to a possible deviation of the salt content values (Figs. 4.15 and 4.16), which were estimated by means of the average HMC of the control-samples, of +6.4% or -3.2% (as a percentage of the estimated salt content values). This means that the climatic conditions in the chamber were quite homogenous.

The average HMC of the pure NaCl samples corresponds to an actual RH of 91.9%. This value was obtained, by linear interpolation, from tables expressing the water activity of NaCl solutions at 20°C as a function of their molality (Robinson and Stokes, 2002). As later confirmed, the climatic chamber was actually working around 3% below the set-point of 95% RH. But this fact had no influence on the results because the salt content of the samples was calculated, according to the method proposed in section 3.6, by reference to the actual HMC of the pure NaCl samples and not by reference to thermodynamic data in the literature.

4.4 Results

4.4.1 - Drying of water-filled specimens

The results of the drying experiments performed on specimens DL1 and ML1 with pure water are depicted bellow. Fig. 4.9 presents selected MRI snapshots obtained during each drying experiment. Fig. 4.10 shows the corresponding 1D moisture profiles at cross-sections CS1 and CS2 (Fig. I.10 in Annex I). This type of 1D profile is only presented for pure water.

The two specimens show a similar drying process with pure water, nonetheless a certain time shift between them (Fig. 4.9 and 4.10): (i) the exposed top surfaces remain wet during the first hours; (ii) after around 4 hours these surfaces begin drying faster than the underlying material; (iii) complete drying of the surface seems to occur after around 5 to 10 hours for specimen DL1 and 8 to 18 hours for specimen ML1, as seen in Fig. 4.10.

In spite of this overall similarity, some differences are observed between the two specimens: (i) stone M contains a higher initial amount of moisture than stone D; (ii) drying of stone M is faster than drying of stone D; (iii) at the end of the experiments, stone M is clearly drier than stone D.



Fig. 4.9 – Drying with pure water: MRI snapshots of specimens DL1 and ML1 in side-view (Fig. 4.7). The receding drying front is identified when a dry (dark) surface layer exists on the top of the specimen.



Fig. 4.10 – Drying with pure water: 1D moisture profiles at cross-sections CS1 and CS2 (Fig. I.10 in Annex I) of specimens DL1 and ML1. NMR signal is given in arbitrary units (a.u.). Due to the resolution of the method, the physical limits of the test specimens are estimated to be approximately within the indicated grey vertical line-segments (the drying surface corresponds to the segment on the left).

4.4.2 - Drying in the presence of NaCl

Selected MRI snapshots of the two drying experiments performed with NaCl are depicted in Figs. 4.11 and 4.12. Snapshots of the experiments carried out with pure water on either specimen are also included in these figures to allow accessing the influence of the salt.

As seen in these figures, NaCl does not qualitatively modify the distribution of moisture in the specimens during drying. Indeed, for both specimens, similar image sequences are obtained with pure water and with NaCl solution. Nonetheless, the figures also reveal that: (i) drying is much slower when NaCl is present, as shown by the fact that the images correspond to later moments of drying in the case of NaCl; (ii) contrary to what occurs for pure water, no receding drying front is observed in neither of the NaCl-loaded specimens until the end of the experiments.



Fig. 4.11 – MRI snapshots of specimen DL1 in side-view (Fig. 4.7) during drying with: (a) pure water; (b) 3 molal NaCl solution. For pure water, receding of the drying front is visible already after five hours (the dark area on the top of the specimen DL1 corresponds to the dry surface layer). Similar MRI images are presented for either solution, which correspond, however, to distinct moments of the drying.



Fig. 4.12 – MRI snapshots of specimen ML1 in side-view (Fig. 4.7) during drying with: (a) pure water; (b) 3 molal NaCl solution. Notice that similar MRI images correspond to distinct moments of the drying in the two experiments.

4.4.3 - Overall drying curves

The drying curves presented in Fig. 4.13 were obtained by means of 1D profiles of the type presented in Fig. 4.10. Each point of a drying curve is calculated by adding all the NMR signal values in the two corresponding profiles of the specimen.



Fig. 4.13 – Drying curves

4.4.4 - Evaporation from free liquid surfaces

The results of the evaporation experiments carried out on free surfaces of pure water or saturated NaCl solution are depicted in Fig. 4.14. In this figure, it is seen that: (i) in both environmental conditions the evaporation rate is higher for pure water than for the salt solution; (ii) the faster the evaporation, the lower the ratio between the drying rates of the two liquids; (iii) in neither of the environmental conditions is the evaporation rate of pure water higher than twice the evaporation rate from the NaCl solution.



Fig. 4.14 – Evaporation from free surfaces of pure water and saturated NaCl solution

4.4.5 - Salt deposition

The final salt distribution in specimens DL1 and ML1 is shown in Figs. 4.15 and 4.16 where the grey scale represents the salt content in each layer. It can be seen that in both specimens: (i) a higher concentration of salt occurs in the outer layer of the plaster; (ii) a certain amount of salt is dispersed throughout the entire specimen.

In spite of this, relevant differences are also observed between the two salt distributions: (i) concentration of salt in the surface layer is higher for specimen ML1; (ii) stone M contains less salt than stone D; (iii) in specimen DL1, a salt accumulation peak exists in stone D next to the plaster; (iv) in specimen ML1, a salt accumulation peak is observed in the bedding mortar next to stone M.





Fig. 4.15 - Final salt distribution in specimen DL1. The salt content values are expressed in kg of NaCl by m^3 of open porosity in the material. Both halves of the specimen are in side-view, as depicted in Fig. 4.7.



Fig. 4.16 - Final salt distribution in specimen ML1. The salt content values are expressed in kg of NaCl by m^3 of open porosity in the material. Both halves of the specimen are in side-view, as depicted in Fig. 4.7.

4.5 Discussion

4.5.1 - Salt-induced dampness

The results of the MRI-monitored experiments show that, with pure water (Figs. 4.9 and 4.10), specimens DL1 and ML1 clearly go through a two-stage drying process, despite of their non-homogenous composition. During Stage I, the moisture content in the specimens decreases over time but the moisture distribution remains homogeneous. After around 2h, the exposed surfaces start drying faster. This is a period of transition between Stage I and Stage II, in which a decreasingly significant part of the porosity is involved in transporting the liquid to the surface (Scherer 1990, Rousset-Tournier 2001, for instance). Stage II begins at the total receding of the drying into the material, which occurs at around 6h in specimen DL1 and somewhat latter in specimen ML1 (Fig. 4.10).

With NaCl solution, the distribution of moisture during drying is similar to the case of pure water, as shown by Figs. 4.11 and 4.12. However, because evaporation was slower due to the presence of salt, the liquid inside the specimens has more time to flow towards the surface. As a consequence, Stage I is longer. Indeed, no receding of the drying front is observed until the end of the experiments with NaCl, which lasted 60 hours for specimen DL1 and 35 hours for specimen ML1. These results agree with:

- (a) Theoretical considerations of Hall and Hoff (2002) that argue that an advancement of the evaporation front towards the outer surface is expected to arise from the presence of soluble salts, in comparison to the case of pure water, due to lowering of the vapour pressure gradient.
- (b) Common gravimetric drying experiments performed under different environmental conditions on water-filled materials. These experiments show that a slower evaporation leads to a longer drying Stage I and to the elimination of a higher amount of moisture during Stage I (Rousset-Tournier 2001, for instance).
- (c) Drying experiments carried out at TUE on porous building materials and monitored by a 1D MRI technique (Pel *et al.* 2003, Petković 2005) where slower drying and no receding of the drying front were observed when NaCl was present instead of pure water only.

The present experiments support, therefore, the hypothesis put forward in section 4.1 that salt-induced dampness is a general feature that arises due to lowering of the evaporation rate by the presence of soluble salts, hygroscopicity being a particular case of that general feature.

As explained, hygroscopic absorption occurs when the environmental RH is higher than the RH_{eq} of the contaminant solution. The negative vapour pressure gradient that exists, in this case, between the salt-loaded surface and the environment is what gives rise to water vapour diffusing from the environment to the masonry and, thereby, increasing its moisture content.

This vapour pressure gradient is positive when, oppositely, the environmental RH is lower than the RH_{eq} of the salt solution. No hygroscopic absorption takes place in this case because water vapour diffuses to the environment and, hence, the salt-loaded masonry dries. However, because salts are present, the drying process is slower than it would be for pure water. In drying tests, this effect induces a longer presence of moisture at the surface, as shown by the present MRI-monitored experiments. In building walls, where the liquid supply flux is often reasonably constant over time, this effect corresponds to the drying front being located at the surface for lower liquid supply fluxes. That implies an increase of the damp surface area by comparison to what would happen if only pure water was present. Indeed, it is known that a slower evaporation causes an increase of capillary rise in walls. This indicates that masonry dampness may be aggravated by the presence of soluble salts even when the RH is lower than the RH_{eq} of the contaminant solution, hence, even if no hygroscopic absorption takes place.

This conclusion is consistent with many practical observations that larger moist areas are recurrently found in salt-loaded masonry (Massari and Massari 1993, Burkinshaw and Parret 2004, for example). However, it contradicts the conclusions of Benavente *et al.* (2003). These authors performed gravimetric drying experiments on stone specimens either saturated with pure water or with a 3 molal NaCl solution. Further, they considered that, for both pure water and the salt solutions, the beginning of drying Stage II corresponded to the inflexion point (Fig. 4.6) where the slope of the drying curve sharply decreased. This assumption lead to the conclusion that the presence of salt reduces both the duration of drying Stage I and the total amount of moisture eliminated by (Stage I) surface evaporation, which opposes the results of the present MRI-monitored experiments.

For pure water, the drying rate is indeed approximately constant during Stage I, abruptly drops when Stage II begins and continues to decrease progressively during Stage II (Sherwood 1929, Scherer 1990). This was observed in the experiments performed here with pure water, whose drying curves (Fig. 4.13) resemble ordinary gravimetric drying curves (Fig. 4.6). Yet, for NaCI a different situation occurred. In this case, the drying curves (Fig. 4.13) show slope variations during Stage I, which was still going on at the end of the experiments (Figs. 4.11 and 4.12). This suggests that the drying rate is not necessarily constant during Stage I when salts are present, a hypothesis put forward also by other authors (Hall and Hoff 2002, Huinink and Pel 2003, for instance). Therefore, gravimetric drying curves of salt-loaded materials cannot be interpreted in the same way as for pure water. That explains the discrepancy in the conclusions of Benavente *et al.*

Decrease of the drying rate of salt-loaded materials during Stage I has been attributed to two different causes (Hall and Hoff 2002, Huinink and Pel 2003): (i) as evaporation proceeds, solutions become more and more concentrated at the drying front, hence, their RH_{eq} and thereby the evaporation rate decrease further; (ii) once saturation is attained, the deposition of salt crystals may hinder evaporation.

The results of the present evaporation experiments on free liquid surfaces of pure water or NaCl solution (Fig. 4.14) suggest that the low drying rates of the NaCl-loaded specimens were probably due to reduction of the specific surface of evaporation. Indeed, the faster the evaporation, the closer the evaporation rates of the two liquids. But even for the faster drying

conditions (20° C-15%RH) the ratio between the evaporation rates of pure water and NaCl solution is only of around 1.4. Therefore, the lower evaporation rate of the masonry specimens in the presence of salt (Fig. 4.13) was not only due to the RH_{eq} of the salt solution being lower than that of pure water. Indeed, these drying experiments were performed under environmental conditions (0%RH and 18°C) that are expected to generate a faster evaporation than the two experiments on free liquid surfaces. Hence, the ratio between the (Stage I) evaporation rates of the contaminated and non-contaminated specimens should be even lower than 1.4. Yet, the evaporation rate was always at least 6 times (specimen DL1) and 3 times (specimen ML1) higher for pure water than for the NaCl solution (Fig. 4.14). Therefore, it is clear that some factor other than the lower RH_{eq} of the salt solution made a major contribution towards decreasing evaporation.

Deposition of salt crystals with a blocking effect was possibly a major cause. But further research, namely at the pore scale, is required to confirm this hypothesis. The suggestion of Rousset-Tournier (2001) that the curvature of the menisci at the drying front could influence the effective surface of evaporation in pure water-filled materials also deserves attention. Indeed, the contact angle of salt solutions with porous building materials is higher than that of pure water and, hence, a lower effective surface of evaporation can arise. It would also be interesting to know if and how the presence of solid crystals can affect that contact angle.

4.5.2 - Salt distribution

The final NaCl distribution in specimens DL1 and ML1 was evaluated by means of HMC measurements which provided estimates of the salt content in distinct layers of the specimens (Figs. 4.15 and 4.16).

The salt distributions depicted in Figs. 4.15 and 4.16 show that overall similarities exist between the two specimens:

- i) A higher concentration of salt always occurred at the outer layer of the plaster, clearly because the moisture was eliminated mostly by (Stage I) surface evaporation.
- ii) The second most superficial layer of plaster always presented a high salt content too. This probably means that the drying front progressed inwards into the specimens during the final oven drying at 40°C.
- iii) The deposition of a certain residual amount of salt throughout the whole specimen can be explained by the decrease in the moisture content at the pore level (Rose 1963b) during the final oven drying. Critical moisture content is attained at a certain moment, at which liquid continuity is interrupted in the capillaries. Isolated clusters of liquid remain thus dispersed in the porous system, trapped in the finer pores. Further drying occurs by local evaporation, hence, in the case of a salt solution, a certain amount of salt will crystallize at each cluster.

Nonetheless of these overall similarities, the final salt distribution is not completely identical in the two specimens due to differences in the pore system of stone D and stone M. Table 4.1 and Fig. 4.8 depict the porosity and porometry, respectively, of the three materials in these specimens. It is seen that:

- Porosity and, thus, the moisture storage capability of stone M is higher than that of mortar L, which is higher than that of stone D.
- Mortar L has less medium-dimension pores (between 0.1 and 0.6 μ m) but more coarse pores (larger than 0.6 μ m) and significantly more fine pores (smaller than 0.1 μ m) than stone D.
- Mortar L has less medium-dimension and coarse pores (between 0.2 and 5 μm) but more fine pores (less than 0.2 μm) and also more coarse pores (larger than 5 μm) than stone M.

The main differences between the two salt distributions are the following:

- The concentration of salt in the plaster, especially in the surface layer, is higher for specimen ML1. This is most likely due to the fact that specimen ML1 has a higher porosity and, hence, contains a higher initial amount of salt solution.
- ii) Stone M, despite its higher initial solution content, accumulates less salt than stone D. In fact, a pore can only remove water from another pore if it has higher capillary pressure, that is, if it is smaller. Stone M has a larger amount of coarse pores than stone D (Fig. 4.8) and, thus, mortar L is able to drain stone M faster and more completely. This was clearly observed in the drying experiments with pure water (Figs. 4.9 and 4.10).
- iii) In specimen ML1, a particularly high concentration of salt exists in the bedding mortar in the contact layer with the stone (Fig. 4.16). This is most likely due to the following. Stone M dries probably before the bedding mortar dries, as it happened in the experiments with pure water (Figs. 4.9 and 4.10). Hence, when the stone is already dry, the mortar may still contain enough moisture to ensure liquid continuity in the capillaries. In that case, a transverse drying front develops at the mortar/stone interface and some salt accumulates in the first contiguous mortar layer.
- iv) In specimen DL1, a higher accumulation of salt occurred in the stone, close to where it came into contact with the plaster (Fig. 4.15). This is probably due to the fact that the critical moisture content of porous materials is higher for less homogeneous pore size distributions (Snethlage and Wendler 1997). This is the case with mortar L when compared to either of the stones (Fig. 4.8). Thus, although with pure water stone D is slightly dryer than the adjacent plaster layer at the end of the experiment (Fig. 4.10), liquid continuity may be interrupted first in the plaster. In this case, a drying front develops at the plaster/stone interface. The drying experiments with salt solution ended before this type of behaviour could be observed, therefore, further confirmation is needed.

This conclusion agrees with previous 1D drying experiments on plaster substrate specimens (Petković 2005). It was found that a fine-pored plaster applied over coarse-pored substrate leads to crystallization only at the surface of the plaster. This could be the case here of plaster L and stone M. Differently, a coarse-pored plaster applied over a fine-pored substrate leads to crystallization both at the surface and at the interface plaster/substrate, as it happens here with plaster L and stone D.

These differences between the salt distributions in the two specimens suggest that, in buildings, distinct local distribution patterns may arise for different masonry materials. However, it is not easy to directly extrapolate the present results and conclusions for porous materials in real constructions. In fact, salt deposition patterns depend on multiple and interrelated factors as it will be shown on Chapter 5.

The environmental conditions, for instance, strongly condition the speed of the drying process. The relative drying speed of the different masonry materials conditions the occurrence of internal drying fronts, either at the pore or at the material scale. It is reasonable to expect that, under slower evaporation conditions, all pores and materials in the masonry will have more time to drain the liquid solutions. Hence, less salt will be deposited in the interior of the masonry, either dispersed throughout the materials (particularly, those of less homogenous pore size distribution) or concentrated close to interfaces between distinct materials.

4.5.3 - Other soluble salts

Only NaCl was studied here. Other salts may have different RH_{eq}^{sat} and crystallization properties and, so, distinct influences on drying. The different drying rates, and consequent tendency for a longer or shorter drying Stage I, may be one of the reasons why some salts tend to form efflorescence and others to be deposited deeper inside porous materials. This could account for the higher tendency of sodium chloride, which has a lower RH_{eq}^{sat} than namely sodium sulfate, to form harmless efflorescence.

In buildings subjected to rising damp, moisture rises up in walls until a steady state is eventually established, so that the ground liquid supply is balanced by evaporation (Arnold and Zehnder 1990, for instance). Hence, the evaporation rate conditions the height of capillary rise, thus, the height at which salts crystallize (Arnold 1989). When evaporation is lowered by whatever factor, for instance, use of coverings with low vapour permeability, rise of RH, lowering of temperature or reduction of air (wind) velocity, the height of capillary rise and salt deposition increases. As shown in this chapter, the salts themselves influence the evaporation rate. More work is needed to understand what exactly the influencing factors are. Nevertheless, it is clear that properties of the soluble salts, and perhaps of their crystallization process, that influence drying need to be considered in order to understand why some salts tend to crystallize higher on walls, while others tend to be deposited closer to the ground.

The concentration of salt solutions can be a relevant factor. This hypothesis is consistent with Lewin's observations (1982). A 60 cm-high stone column was subjected to in-continuum partial immersion in NaCl solutions with different concentrations. Lewin reports that the height at which damage occurs increases with the concentration of the salt solution, a behaviour that is compatible with the results of this thesis. Also in the experiment performed by Van Hees and Lubelli (2002) on three brick masonry walls, the higher the concentration of the NaCl solution the higher the obtained level of capillary rise. The minimum level was achieved with pure water.

In fact, the RH_{eq} of salt solutions varies (decreases) with their concentration. Therefore, concentration influences the evaporation rate, at least, before saturation is attained. Concentration conditions also the rate of crystallization in porous materials. This may be relevant, as far as evaporation is concerned, if it is true that salt crystals hinder evaporation. Therefore, the concentration of salt solutions may condition masonry drying and salt distribution although the possible relevance of this influence requires further evaluation. Nevertheless, it is reasonable to expect that the lower the concentration, the more similar to the case of pure water is the drying process, i.e., the deeper and higher in walls the drying front is positioned. This would explain why weak salt solutions are able to cause significant damage in old constructions.

NaCl did not seem to significantly affect the interaction between the different materials during the present drying experiments (Figs. 4.11 and 4.12). Although the duration of these experiments was insufficient for a complete comparison of the two drying processes, the final salt distributions are compatible with this hypothesis, as discussed in the previous section. It is necessary to verify, in future work, if the same happens with other soluble salts. Different interaction between the various masonry materials during drying may lead to distinct local salt distribution patterns.

Chapter 5 – Behaviour of plasters and renders

5.1 Introduction

The behaviour of porous building materials in relation to salt crystallization can be influenced by several factors. The environmental conditions, for example, are a major influencing factor. They condition, for instance, the evaporation and crystallization rates, the nature and habits of the crystalline phases that arise or the occurrence of temperature-induced crystallization, as discussed in Chapter 2.

The type of salt is another important factor. It is known that different salts are effective under different environmental conditions (Goudie and Viles 1997). Yet, some salts tend to be more harmful than others. In laboratory tests, for instance, sodium sulfate typically causes extensive damage to porous building materials, while sodium chloride is often much less damaging or even completely innocuous. Since these are the two salts normally used in laboratory crystallization tests, it is important to understand the reasons for such different behaviour.

One plausible cause was put forward in Chapter 4: sodium chloride induces a slower drying, namely because of its lower RH_{eq}^{sat} , which could account for a higher tendency to form harmless efflorescence than sodium sulfate.

Despite this apparently low destructive capability, sodium chloride is often found to be involved in salt decay processes in real constructions. This inconsistency between the in-lab ineffectiveness and on-field effectiveness of sodium chloride was pointed out by authors such as Honeyborne (Evans 1969-1970) or Goudie and Viles (1997) and requires further explanation.

Plasters and renders with different working principles are used today on salt-loaded walls. Some are industrial products aimed specifically at being salt-accumulating or salt-transporting. But real buildings present a wide range of possibilities as regards not only the environmental conditions and type of salt but also, for instance, the nature of the substrate, the moistness conditions or the presence of a final paint layer. Therefore, a question arises as to whether and how the behaviour of plasters and renders in relation to salt crystallization can be influenced by such conditions. In Portugal, traditional (site-made) plasters and renders are widely used on salt-loaded walls. Water-repellent mass additives are often used in such plasters and renders aiming at minimizing salt dampness and surface damage by salt crystallization.

It is known that fully hydrophobic surface layers can worsen salt decay because crystallization takes place at the interface between the hydrophobic layer and the non-hydrophobic material. This was verified by the author within a set of NaCl crystallization tests on brick specimens, either non-treated or treated with a surface hydrophobic product (Gonçalves 2003b). After several cycles of partial immersion in a saturated solution of NaCl and oven drying at 60°C, the treated specimen cracked due to subflorescence formation at the interface between hydrophobic and non-hydrophobic material, as seen in Fig. 5.1. But the non-treated specimen presented only efflorescence.



Fig. 5.1 - Crystallization test on brick treated with hydrophobic surface treatment on all faces except the bottom (specimen I) and on non-treated brick (specimen II): (a) damage at the end of the test; (b) damage after removal of efflorescence; (c) subflorescence in the treated specimen, after removal of the fractured surface layer.

Fully hydrophobic plasters and renders, either traditional or industrial, are sometimes used on salt-loaded walls. Their behaviour is expected to be equivalent to that of specimen I in Fig. 5.1, that is, occurrence of subflorescence at the plaster/substrate interface (Fig. 2.10) and ultimate detachment from the substrate. But partially hydrophobic plasters and renders, where low amounts of mass water-repellent additives are used to reduce the capillarity coefficient, are probably the most common situation. Also in this case, there is field evidence that salt decay may be worsened, as observed by the author on test panels applied on an old church near Lisbon (Fig. 5.2). The reasons for salt damage increasing when partially hydrophobic plasters and renders are used required clarification.

Dampness and decay problems are in fact often aggravated by restoration interventions. This is a general complaint from many end-users, as reported in Chapter 2. Such problems are perhaps related to the use of non-traditional products, as in the case presented in Fig. 5.2 which concerns the use of hydrophobic additives. Failures of industrial salt-accumulating plasters have been also reported, for instance, in the case of Leiderdorp Farmhouse in the Netherlands (TNO 2004). It is, therefore, critical to identify what factors can worsen dampness and decay in recently restored buildings.



Fig. 5.2 – Test panels on the interior of S. Sebastião church, Almada, one year after their application. Each panel is made of two similar plaster coats composed by a mortar with volumetric composition of 1:1.5:1.5 (dry-hydrated lime: sand from Tagus river : yellow pit sand from Corroios). In the panel on the right a commercial air-lime with hydrophobic properties was used. In contrast to the panel on the left, where ordinary air-lime was used, more severe dampness and sanding of the mortar at the base of the panel were observed here.

Laboratory tests were considered the best means of answering the previous questions. Because of their operational advantages, laboratory tests are, in contrast to field tests, a feasible means of evaluating many types of materials under different conditions. Further, laboratory tests can be performed under controlled conditions, hence, allowing sustainable comparisons. Laboratory tests may also be designed in such a way as to isolate possible influencing factors.

Laboratory salt crystallization tests are currently used to assess or predict the performance of building materials. For plasters and renders, though, there are no standard crystallization tests available. In fact, there were no available methods of whatever kind allowing an adequate evaluation of the performance of plasters and renders on salt-loaded walls. Yet, the influence of plasters and renders on moisture transport and salt crystallization processes throughout an entire building can be huge, as sustained in this thesis. Further, more and more industrial products, of varied nature and often considerably expensive, which are claimed to prevent salt-induced dampness or salt decay problems, are available on the market. Hence, there is an urgent need for methods that will allow a preliminary evaluation of the behaviour of plasters and renders for salt-loaded walls.

For all these reasons, a laboratory research was carried out in order to investigate the behaviour of plasters and renders in relation to salt crystallization. The main objectives were:

- To assess the behaviour of three different plasters, either painted or unpainted, with different salts and substrate materials. Two commercial plasters, intended to work as salt-accumulating, and a traditional lime-cement plaster were used.
- To evaluate the possible influence of the following main factors:
 - moistness condition;
 - type of salt;
 - substrate material;
 - presence of a paint layer;
 - use of water-repellent additives in traditional plasters and renders.
- To identify chief factors that can account for the worsening of dampness and salt decay after restoration interventions.
- To assess the possibilities and limitations of a given crystallization test as a performance test for plasters and renders.
- To discuss the adequacy of plasters and renders with different working principles to different practical situations.

The research was performed by means of salt crystallization tests or MRI-monitored drying experiments. Five sets of tests were performed in total. The work was done within different research programmes in which the author was involved over recent years. For this reason, the choice of the testing procedures and materials was to a large extent conditioned by the specific requirements of each programme.

In this chapter, the overall research is presented. Salt crystallization tests are reviewed in section 5.2. Background information on the sorption behaviour of porous building materials is introduced in section 5.3. The materials used in the five sets of tests are described and characterized in section 5.4. Then, the results of each set of tests are individually presented and discussed in sections 5.5 to 5.9. An overall discussion focusing on the objectives of the research is presented in section 5.10.

5.2 Salt crystallization tests

5.2.1 - Introduction

Salt crystallization tests are ultimately drying tests performed on specimens filled with a salt solution. Still, there are important differences between salt crystallization tests and ordinary drying tests such as those described in Chapter 4. One is that salt crystallization tests are not normally performed under constant conditions. They often involve periodic wetting and drying under changeable environmental conditions. But the fundamental difference is that salt crystallization tests are aimed at evaluating salt damage and not only moisture transport during drying. Salt crystallization tests are in fact artificial ageing tests. Very concentrated salt solutions, repeated contaminations and severe or changeable environmental conditions are used in order to enhance damage by inducing fast drying and crystallization rates, increased crystallization pressures, temperature-induced crystallization, hydration, etc.

Typically, in salt crystallization tests the specimens absorb the chosen salt solution under a predefined contamination procedure. Afterwards they are allowed to dry under certain environmental conditions. Often, several wet/dry cycles are carried out. Differently, some crystallization tests are performed on specimens in continuous partial immersion. In this case, absorption and drying occur simultaneously throughout the entire testing period. In both types of test, damage is evaluated, either periodically during the test or at the end, by visual observation or by measuring certain material properties.

The first crystallization tests were carried out in the 19th century by civil engineers for simulating the disruptive action of icing. In his early article, Thury (1828) reports the method proposed by M. Brard: small cubes of stone were boiled for 30 minutes in a sodium sulfate solution saturated when cold; the cubes were then hung above dishes containing the same salt solution and were periodically sprinkled with water for around five to six days; the resistance of the stones to icing was evaluated by considering the loss of material.

Salt crystallization itself was later recognized as an important deterioration mechanism of porous materials. So, crystallization experiments began being performed to specifically assess damage by salt crystallization. Schaffer (1932), for instance, states that crystallization tests: (i) allow correlating significant factors, namely environmental conditions and material structural properties; (ii) tend to reproduce that type of weathering which the material suffers under actual exposure to the weather.

Eventually, some crystallization tests were included in official standards, for example, CEN test for natural stone (CEN 1999) or ASTM test for aggregates (ASTM 2005). But, no crystallization tests specific for plasters/renders neither exist as standard methods nor are mentioned in article or book reviews on the subject (Evans 1969-1970, Alonso *et al.* 1987, Goudie and Villes 1997).

In the late nineties, Wijfels *et al.* (1997) proposed a crystallization test for plasters/renders. This test is similar to a crystallization test for masonry wallets by RILEM, latter published as RILEM (1998), with the following main alterations: (i) it was performed on specimens composed of a plaster on brick substrate; (ii) at the end of the test, the salt distribution across the specimens was measured.

More recently, efforts have been made (Vergès-Belmin *et al.* 2005) within the research project COMPASS to develop an effective crystallization test for plasters and renders by improving the method used by Wijfels *et al.* Some of the experimental work presented in this Chapter (section 5.5) was part of that common effort.

5.2.2 - Test variables

Alonso *et al.* (1987) identified some of the main variables of crystallization tests by wet/dry cycles which are the most common case:

- specimens: size, shape and number;
- salt: nature of the salt and solution concentration;
- contamination phase: type of immersion (total or partial), duration and temperature of the solution;
- drying phase: environmental conditions and duration;
- cooling phase: existence and duration;
- number of absorption/drying cycles;
- damage assessment: by visual observation, by considering the number of cycles the material resists, by mass loss evaluation or by accessing changes in physical properties such as, for instance, porosity.

Other test variables, relevant for crystallization tests in general and not only for those by wet/dry cycles, are:

- type of specimens: single-material or composite specimens;
- general procedure: wet/dry cycles, single initial contamination or continuous immersion of the specimens;
- contamination procedure: in the case of wet/dry cycles or single initial contamination, the specimens are contaminated by allowing them to absorb the solution during a given period; this period is either equal for different types of materials or predetermined in order to correspond to the introduction of similar amounts of salt or salt solution in these materials; in the case of wet/dry cycles, further rewettings can be carried out with the same salt solution or with pure water only.

- drying procedure: unidirectional migration of moisture (by sealing the lateral faces of the specimens) or not;
- assessment of damage: periodicity; measurement or not of the final salt distribution;
- stopping criterion: accomplishment of a certain number of cycles or occurrence of damage.

The most significant of these test variables will be discussed in the following sub-sections in the light of an adequate design of crystallization tests for plasters and renders.

5.2.3 - Type of specimens

Most crystallization tests reported in the literature are meant to assess the resistance/durability of certain material(s) to salt crystallization. Accordingly, they are performed on specimens made of a single material (concrete, mortar, brick, stone, etc.). However, for plasters and renders, an evaluation of the material itself is clearly insufficient:

- a) Firstly, because the physical properties of plasters and renders may be influenced, during hardening, by those of the substrate material.
- b) Secondly, because solution transport, hence, salt crystallization and salt distribution processes depend on the interaction between all masonry materials (Huinink *et al.* 2005, for instance).
- c) Thirdly, because plasters and renders do not work independently of the substrate: their detachment is already damage, even if no other deterioration occurs in the plaster/render or in the substrate materials. Fully hydrophobic plasters, for instance, will certainly perform very well if single-material specimens are used because they do not absorb liquids. However, in reality, fully hydrophobic plasters and renders often cause problems, namely, when they detach from the masonry.
- d) Finally, because plasters and renders have a fundamental sacrificial role in ancient buildings: they are meant to protect the structural masonry from destructive actions, namely, salt crystallization. Although in some cases other requirements, such as providing good health conditions inside buildings, may superimpose on substrate protection requirements, it is therefore important to verify to what extent plasters and renders prevent or, conversely, encourage salt crystallization in the substrate.

RILEM test MS-A.1 (RILEM 1998) is one of the few crystallization tests that use composite specimens. These are the so-called wallets which are composed of a number of masonry elements and mortar joints. The large size of these wallets (minimum flat dimensions of 200 mm x 200 mm) seems, however, to be a significant operational disadvantage.

Although the use of plaster/render is not foreseen in RILEM recommendations, nothing in the test method itself prevents its use. Wijfels *et al.* (1997) in fact performed RILEM test MS-A.1 on specimens composed of plaster applied on brick substrate (apparently, entire bricks of unspecified dimensions were used).

A recent NaCl crystallization test proposed by Lubelli (2006) and the MRI-monitored drying experiments carried out by Petković (2005) also used plaster/substrate specimens. These works were, to a large extent, related to the COMPASS project under which framework this thesis was partially carried out.

5.2.4 - Contamination conditions

In crystallization tests by wet/dry cycles, the specimens are filled with the salt solution generally by one of the following type of protocols: (i) capillary saturation by total immersion; (ii) introduction of a fixed amount of salt, by keeping the specimens in partial immersion during a predetermined period, which can vary for different materials; (ii) partial immersion of all types of specimens during a fixed period.

Capillary saturation can perhaps be representative of reality for very absorbent materials or, ultimately, materials in walls where a moisture supply rate higher than the evaporation rate exists during very long periods. But it may be unsuitable for simulating the condition of plasters applied on walls where wetting and drying periods alternate. In this case, plasters with high vapour permeability and low capillary absorption will probably never attain capillary saturation, while very absorbent plasters may do so. Therefore, capillary saturation can be an erroneous condition when these two kinds of plasters are compared.

RILEM test MS-A.1 (RILEM 1998) uses a multi-stage contamination regime. The wallets are immersed in the solution up to 20 mm from their bottom face. Four hours later that level is restored. After another four-hour period, the solution level is brought up to 5 mm below the top surface of the wallets. The specimens are removed from the solution when 24 h have passed since the start of contamination. This multi-staged contamination procedure is mostly aimed at avoiding air-trapping but, excepting that, it is practically equivalent to capillary saturation by total immersion.

Representativeness of the second condition (fixed amount of salt) is also questionable, either when it corresponds to similar amounts of salt or to similar salt contents in different specimens, because different building materials in similar field conditions are likely to absorb distinct amounts of solution. Indeed, introducing a given amount of salt can be impossible, for example, in the case of hydrophobic materials.

The use of a fixed amount of salt may, nevertheless, be appropriate in certain cases. For example, Lubelli (2006) used identical NaCl contents of 2% (by weight) in traditional plasters in order to reproduce the salt content in certain masonry.

The third possibility, partial immersion during a fixed period, may not bring about these nonrepresentativeness disadvantages. In that case, different plasters are subjected to an action of similar intensity. Like in reality, the more absorbent plasters will absorb a higher amount of solution. However, evaluating the behaviour of specimens that contain very different amounts of salt may be difficult, particularly when salt distribution profiles are to be compared.

In some cases, an initial contamination with salt solution and further re-wettings with only pure water are carried out. This is expected to be a less damaging condition than when successive contaminations with the same salt solution are carried out. This procedure was used on the preliminary crystallization tests in Fig. 5.3, although it is hardly sustainable as representing reality.

Perhaps more interesting are the procedures where a single contamination is carried out at the beginning of the test, typically by total immersion in the salt solution (Faria Rodrigues 2003, for instance). It has been argued that these so-called non-immersion techniques are probably more representative of reality (Goudie 1997).

Procedures based on continuous (partial) immersion of the specimens are sometimes used instead of wet/dry cycling, although they are less common. This type of procedure was evaluated in the preliminary crystallization tests in set 1, as described in section 5.5.2.1 and depicted in Fig. 5.9.

5.2.5 - Drying procedure

In real walls, salt solutions typically migrate from the interior of the masonry towards the outer surface of the plaster/render. This situation is normally represented in laboratory tests by sealing all the faces of the specimens, except that representing the outer surface of the wall. This prevents lateral evaporation during drying, thereby enabling predominantly unidirectional migration of moisture to the surface.

The situation is not so straightforward as regards the environmental conditions during drying. Enhanced damage is ordinarily achieved in crystallization tests by means of high temperatures or low RH. But a large range of conditions is used in practice.

Drying is sometimes performed under constant environmental conditions of temperature and RH. This happens, for instance, in RILEM test MS-A.1 (RILEM 1998) which is carried out at 20°C and 50%RH. In other cases, oven drying at high temperatures is performed cyclically between contaminations. This is the case of ASTM C88 test (ASTM 2005), which uses $110\pm5^{\circ}$ C or RILEM test V.1.a (RILEM 1980) where a temperature of $105\pm5^{\circ}$ C is to be reached after 10 to 15 hours. Oven drying at a lower temperature of 60°C is used, for example in RILEM tests V.1.b or V.2 (RILEM 1980), to prevent alteration of surface treatments.

Drying, particularly in non-standard salt crystallization tests, often includes RH or temperature cycles. The aim is sometimes to represent specific environmental conditions as, for instance, in Goudie's climatic simulations (Goudie and Viles 1997). More often, environmental cycling is intended to cause cycles of hydration/dehydration, deliquescence/crystallization or temperature-induced crystallization.

In some non-immersion techniques the specimens are oven dried after being contaminated and before being exposed to the testing conditions. This happens, for instance, in the crystallization tests reported by Faria Rodrigues (2003), which use cycles of 12 h at 90%RH and 20°C and 12h at 40%RH and 20°C.

When both hygroscopic moistening and drying occur during the exposure period of the specimen, typically in alternating phases of high and low RH, it is not totally adequate to designate that period as "drying phase". This happens in non-immersion tests, which are ordinarily based on hygroscopic moistening, but also in some wet/dry cycle tests that include RH cycling.

5.2.6 - Salt solutions

Very concentrated or even saturated solutions, typically of sodium sulfate or sodium chloride, are generally used in crystallization tests. Since sodium sulfate and sodium chloride are also amongst the most common salts in decayed buildings and, moreover, were the ones used in the experiments of this thesis, the following discussion will focus on them.

Different salts are effective under different environmental conditions (Goudie and Viles 1997). Severe damage is normally achieved with sodium sulfate in crystallization experiments, which is probably why this is the salt most often used in official standard methods (Alonso *et al.* 1987). In contrast, sodium chloride, although frequently involved in salt decay processes in buildings, is often found to be one of the least damaging or even a totally innocuous salt in laboratory tests.

Sodium sulfate was already chosen by Brard for his early crystallization test on stone (Thury 1828) for being more destructive than sodium chloride, potassium nitrate, magnesium sulfate and sodium carbonate. The environmental simulations with varying temperature and RH carried out by Goudie on concrete specimens (Goudie and Viles 1997) are a more recent example. Sodium sulfate was the most effective of five salts in one of the six conditions (although it was ineffective under the five other conditions), while sodium chloride was the only salt that had no effect under any of the conditions.

Within a preliminary set of crystallization tests in this thesis, more severe damage was also achieved with sodium sulfate than with sodium chloride, except in one of the tested environmental conditions (Fig. 5.3).

	20°C - 50% HR (conditioned room, low air velocity)	50°C - 50% HR (climatic chamber, low air velocity)	20°C - 35% HR (climatic chamber, low air velocity)	1 week: 20°C-50% RH (conditioned room, low air velocity) 1 week: 20°C-≈90% RH (container, over water)	≈22°C - 70 to 75% RH (non-conditioned room, despicable air velocity)
NaCl 1.9 molal (10%W)	Efflorescence+sanding	Efflorescence+sanding	Only efflorescence developed very tight to the surface of the plaster.	Only efflorescence (whitened surface) during the low RH periods	Only powdery efflorescence
	55 days (2 rewet)	42 days (6 rewet)	45 days (4 rewet)	38 days (no rewet)	30 days (no rewet)
Na₂SO₄ 0.39 molal (5%W)	Exfoliation	Neither efflorescence nor any other visible type of damage	Scaling	Bulging	Efflorescence+sanding
	14 days (1 rewet)	55 days (7 rewet)	16 days (2 rewet)	28 days (1 rewet)	2 days (no rewet)

Fig. 5.3 – Preliminary crystallization tests by wet-dry cycles. The specimens are composed of a 15 mm thick air-lime plaster applied on 2 0mm thick brick substrate and have flat dimensions of 20 mm x 20 mm. They are contaminated with NaCl or Na_2SO_4 by absorbing the solutions through their bottom faces during 20 min, which corresponds approximately to reaching 80% of the maximum capillary moisture content. Then, the specimens are subjected to the reported environmental conditions. They are rewetted with pure water only at the end of each drying period, whenever their mass is approximately constant. The test carried out in a non-conditioned room (column on the right) is an additional test performed on saturated (undamaged) specimens at the end of the continuous immersion tests presented in section 5.5.2.1. These preliminary crystallization tests ended at the observation of damage other than efflorescence or when damage was no more expected to occur within a reasonable period.

Rodriguez-Navarro and Doehne (1999) also obtained less damage from sodium chloride than from sodium sulfate. Their crystallization experiments on stone blocks were performed by continuous partial immersion under constant environmental conditions at a low (35%) or medium-high (60%) RH. Though both salts induced more damage under low RH conditions, sodium sulfate formed mostly subflorescence in highly localized areas, while sodium chloride predominantly formed efflorescence or filled the smallest pores homogeneously.

5.2.7 - Damage assessment

The performance of plasters and renders in relation to salt crystallization can be evaluated as regards the damage patterns that occur, the durability or the salt accumulation behaviour of the plaster/render.

Damage patterns are normally evaluated by visual observation. In some cases visual observation is complemented, for instance, with microscopy (Alonso *et al.* 1987).

Durability is typically assessed by using quantitative parameters such as the number of cycles accomplished (or time passed) before damage is observed or the mass loss at the end of the test or at the end of each cycle.

The salt accumulation behaviour can be disclosed by the final salt distribution across the plaster/render. Though it is not measured in ordinary salt crystallization tests, the salt distribution was evaluated, for instance, at the end of the tests performed by Wijfels *et al.* (1997) on brick/plaster specimens. First, the specimens were sliced. Afterwards, the 3 mm slices obtained were dried, weighed and stored at 20°C and 93% RH (for an unspecified period) over a saturated sodium sulfate solution. Relative evaluation of the salt content in the slices was made by means of their HMC. Ion chromatography is sometimes used instead of the HMC method to obtain distribution profiles, for instance, in the experiments carried out by Petković (2005).

5.3 Sorption of porous building materials

The sorptivity of porous building materials varies with their moisture content. This happens because, when a material is wet, a certain part of its porosity is saturated and, hence, does not contribute to the overall suction capability.

It is a common assumption that, at given moisture content in a material, all the pores below certain dimensions are saturated. However, in reality, the distribution of water in the complex pore networks of porous building materials can be significantly different when it is reached by absorption or desorption. Several factors have been argued to contribute to this difference, namely, pore shape, pore wall surface roughness or air-trapping during absorption (Bachmann and van der Ploeg 2002). Nevertheless of the cause, when pores of different dimensions are saturated, the capillary suction of the material is different. Therefore, the material sorptivity can depend on whether the moisture content was achieved by wetting or drying the material, which configures a hysteretic effect.

Water characteristic curves (Fig. 5.4) express the correlation between the moisture content in a material and its matric potential which represents the energy necessary to remove a unit weight of water from the porous solid to a free state at the same level. Since in porous building materials capillary forces are predominant (except at very low water contents where adsorption is relevant) the matric potential can be, for practical purposes, assumed to be equal to the capillary potential.



Fig. 5.4 – Water characteristic curve of a porous building material

The capillary potential ψ (m) is given by Eq. 5.1, where p_c (Pa) is the capillary pressure, ρ (kg.m⁻³) the bulk density of water and g (m.s⁻²) the gravitational constant.

$$\Psi = \frac{p_c}{\rho.g}$$
 Eq. 5.1

Due to the hysteretic nature of the capillary potential, water characteristic curves (Fig. 5.4) have two branches: one corresponding to absorption (wetting branch) and the other to desorption processes (drying branch).

The equilibrium condition for two different wet materials in contact is the equality of capillary potential. This does not normally correspond to an equality of moisture content because of the differences between the two pore networks.

The liquid flux in porous building materials can be expressed in terms of capillary potential, as shown by Eq. 5.2. that concerns unidirectional migration of moisture.

$$u = K_1(\theta) \frac{\partial \Psi}{\partial l}$$
 Eq. 5.2

In this equation, u (m³.m⁻².s⁻¹) is the liquid flux, K_1 (m.s⁻¹) the liquid conductivity of the plaster which is a very strong function of the moisture content and I (m) the distance.

5.4 Material characterization

Table 5.1 presents the materials on which all the experiments presented in this chapter were carried out.

The mortars, brick and stones indicated in Table 5.1 were characterized by means of the following tests:

- capillary absorption tests were performed (Fig. 5.5) on all mortars, brick and stones used in the five sets of tests;
- drying tests were carried out (Fig. 5.6) on the mortars used in sets 1 and 2;
- porosity and pore size distribution measurements (Fig. 5.7) by mercury intrusion porosimetry (MIP) were performed on the mortars, brick and stones of sets 2 to 5;
- water vapour permeability tests (Fig. 5.8) were carried out on the three plastering mortars used in set 1; these tests were performed by IET within the COMPASS project and kindly made available for this thesis.

Designation	Composition / description		
Mortar L	Traditional mortar with volumetric composition of 1:1.5:1.5 (dry hydrated lime LUSICAL : sand from the Tagus river : yellow pit sand from Corroios)		
Mortar H	Similar to mortar L but including a mass hydrophobic agent that was added during batching in the proportion of 1/50 (by weight) to the lime		
MEP-SP®	Industrial trass-(air)lime-based plastering/rendering mortar by Strasserville (France), which includes mass water-repellent additives		
Parlumière®	Industrial cement-based plastering/rendering mortar by Lafarge Mortiers (France), which includes mass water-repellent additives	1	
LNEC-plaster	Traditional (air)lime-cement-based plaster/render composed of an adherence coat and a base coat with volumetric composition of, respectively, 1:1:6 and 1:3:12 (type IV cement : dry-hydrated lime LUSICAL : river Tagus sand)	1	
Brick	Dutch red ceramic brick (solid bricks)	1 and 2	
Stone D	Portuguese medium-porosity calcitic calcareous stone with fossil fragments (currently named "vidraço")	3, 4 and 5	
Stone M	Portuguese high-porosity calcitic calcareous stone	4	
Silicone paint	Funcosil® LA silicone-based emulsion paint (ochre) by Remmers	1	
Acrylic paint	DIOPLASTE® average-quality acrylic emulsion paint (white) by Barbot (Portugal)	3 and 5	
Mass hydrophobic additive	SUPER SIKALITE® mass hydrophobic additive (powder) for mortar and concrete by Sika (Portugal)	4	

Table 5.1 - Materials used in the specimens

The unidirectional capillary absorption tests (Fig. 5.5) were carried out on specimens with dimensions of around 45mm x 45mm x 20mm whose lateral faces were sealed with an epoxy resin. Only in the case of lime mortars L and H was the test carried out according to CEN's procedure (CEN 2002) on six half-prisms, laterally sealed with wax, obtained by breaking three 40 mm x 40 mm x 160 mm prisms. In both cases, the specimens were first oven dried at 60°C. Then, they were allowed to freely absorb pure water through their bottom face and were weighed periodically. During absorption the specimens were kept inside a closed acrylic box and the level of the free water surface was maintained 5 mm above the bottom face of the specimen. The test was performed on two or three specimens of each material. The water absorption coefficient (WAC) is calculated as the inclination of the straight (initial) section of the absorption curve which expresses the average weight amount of water absorbed per unit area as a function of the square root of time.



Fig. 5.5 - Capillary absorption tests (WAC in kg.m⁻².h^{-1/2}).

The drying tests were carried out in a conditioned room with a low air velocity at approximately 20°C and 65% RH. Plaster slabs of around 45 mm x 45 mm x 20 mm, laterally sealed with an epoxy resin, were used. The slabs were detached from plaster/brick specimens similar to those in Fig. 5.10 but where filter paper was interposed between the plaster and the substrate. They were previously saturated by total immersion in pure water until constant weight. During drying, their bottom face was sealed with polyethylene film. Three specimens of each kind were used.



Fig. 5.6 - Drying curves

The pore size distribution was measured by mercury intrusion porosimetry (MIP) according to LNEC's procedure LERO PE01, which is based on NP EN 1936 (CEN-IPQ 2001).



Fig. 5.7 – Pore size distribution (MIP)

The material vapour transport capability was characterized by means of RILEM test II.2 (RILEM 1980). The tests were performed on plaster slabs detached from plaster/brick specimens, as in the case of the drying tests presented in Fig. 5.6.



Fig. 5.8 - Water vapour transport. Results obtained and kindly made available by IET.

5.5 Crystallization tests on plaster/substrate specimens (set 1)

5.5.1 - Introduction

Salt crystallization tests were performed on three different plasters applied on brick or on lime mortar substrate. Two commercial plasters claimed to work as salt-accumulating plasters and a traditional lime-cement plaster were used. Tests with either sodium chloride or sodium sulfate were carried out on painted and unpainted specimens.

The main objectives of these tests were the following:

- Assessment and comparison of the behaviour of the plasters, particularly as regards their salt-accumulation behaviour, under different conditions;
- Evaluation of the influence of the following main factors:
 - type of salt, by comparing the results obtained with NaCl or Na₂SO₄;
 - substrate material, by comparing the performance of the traditional plaster on brick or on lime mortar substrate;
 - presence of a paint layer, by comparing the results obtained with painted and unpainted specimens;
- Discussion of the testing protocol.

The tests presented in this section were carried out at LNEC within the COMPASS project (Vergés-Belmin *et al.* 2005).

5.5.2 - Experimental conditions

5.5.2.1 - Definition of the main testing protocol

The COMPASS salt crystallization test derives from the test reported by Wijfels *et al.* (1997) which is carried out on plaster/brick specimens. The test of Wijfels *et al.* is based on RILEM test for masonry wallets (RILEM 1998) as regards, namely, the contamination protocol and environmental conditions during drying. The COMPASS test, however, includes changes which are mostly aimed at increasing its effectiveness. Indeed, the conditions of RILEM test can lead to a very slow damage velocity. For instance, in previous tests carried out according to this protocol on brick wallets with different surface treatments (van Hees and Brocken 2004), most damage occurred approximately between 12 and 48 months, which is hardly an acceptable period for most purposes.

For the present experiments, two methods were initially considered: (i) wet-dry cycles including an initial contamination with the salt solution and further rewettings with pure water only; (ii) continuous partial immersion of the specimens in the salt solution. The possibilities of these methods were evaluated by means of preliminary tests performed on plaster/brick specimens under different environmental conditions. A pure lime plaster was used at LNEC. An industrial restoration plaster was tested in two other laboratories (TNO and LRMH). The final salt distribution was obtained, in the case of the industrial plaster, by slicing the specimens and measuring the HMC of each sample. The different environmental conditions used in these preliminary tests are shown in Table 5.2 where those used in LNEC tests are marked.

Test by	Reference 20°C - 50% HR	High T 50°C - 50% HR	Low RH 20°C - 35% HR	Cycles of RH 1 week cycles 20°C - 50% RH 20°C - ≈90% RH	Cycles of T 1 week cycles 20°C - 50% RH 50°C - 50% RH	Cycles of T and HR
Wet-dry cycles	х	х	х	х	-	-
Continuous immersion	х	x	х	-	Х	-

Table 5.2 – Environmental conditions used in the preliminary crystallization tests

20°C/50% RH - conditioned room, low air velocity; 50°C/50% RH - climatic chamber, low air velocity;

20°C/35% RH - climatic chamber, low air velocity; 20°C/≈90% RH - over water in a closed container at a conditioned room

The results of the preliminary crystallization tests carried out at LNEC by wet-dry cycles or continuous-immersion are shown in Figs. 5.3 and 5.9, respectively. Fig. 5.3 provides also details on the testing conditions. As seen, an additional test was performed at LNEC with no rewettings and drying in a non-conditioned room.



Fig. 5.9 - Preliminary crystallization tests by continuous immersion on lime-plaster/brick specimens. Abundant efflorescence occurred under all environmental conditions both for the 1.9 molal sodium chloride (on the left) and the 0.39 molal sodium sulfate (on the centre) solutions. Its removal systematically revealed wet undamaged plaster surfaces. The only exception were the tests with high temperature periods (constant or cycles) on sodium sulfate, where cracking of the specimens was observed (on the right).

The preliminary tests performed at TNO and LRMH on the industrial plaster allowed concluding that wet-dry cycles induced higher salt contents in the plaster than continuous immersion (TNO 2003). The tests performed at LNEC on the traditional plaster showed that that continuous immersion was in general a much less harmful or even totally ineffective method (Fig. 5.9) in comparison to wet-dry cycling (Fig. 5.3). Therefore, wet-dry cycles were the method chosen for further tests.

Another conclusion was that high temperature originated faster drying, hence, more rewettings in the case of wet-dry cycles. For this reason, it was decided to use a drying period at a high temperature. In order to accelerate drying without using a very high, thus perhaps too unrealistic, temperature, it was decided to combine this high temperature with a low RH. The chosen conditions were 40°C and 20% RH. A period at ordinary room conditions was also desirable so that the specimens could be periodically observed and manipulated with minimum disturbance of the testing conditions. Therefore, daily cycles of 16 hours at 40°C and 20% RH followed by 8 hours at 20°C and 50% RH were finally adopted.
In the preliminary tests, no significant damage to the industrial plaster arose in none of the conditions (TNO 2003). For this reason, it was decided to harsh the testing conditions by performing rewetting with the same salt solution instead of pure water only.

Small size specimens with flat dimensions of around 5 cm x 5 cm composed of a plaster applied on a homogenous brick substrate were used in the preliminary crystallization tests rather than the large size wallets (25 cm x 20 cm) of RILEM test (RILEM 1998). These small sized specimens showed to be sufficiently representative and, hence, were adopted for the subsequent tests. Indeed, the use of small specimens provides enormous operational advantages: a much higher number of specimens can be tested simultaneously in a climatic chamber, they can be handled faster and more easily and ordinary high precision weighing devices can be used.

A contamination procedure different from that of the preliminary tests (Fig. 5.3) was, however, used in the subsequent tests. This new procedure was aimed at overcoming the disadvantages of common contamination procedures (section 5.2.5), that is, introducing either very different amounts of salt in different specimens or forcing the specimens to absorb equal quantities of solution which can be unrealistic for some materials. In the present case, it was decided to introduce a fixed amount of solution but only in the substrate. Therefore, different plasters could be suitably compared because:

- the substrate contained in all cases the same amount of salt solution, which is representative of a situation where all plasters are subjected to an action of similar intensity;
- identical amounts of salt could be introduced in the specimens regardless of the physical properties of the plaster.

5.5.2.2 - Materials and methods

The present crystallization tests were carried out on industrial plasters MEP-SP and Parlumière and traditional plaster LNEC (Table 5.1). MEP-SP and Parlumière were applied only on brick substrate. LNEC was applied either on brick or lime mortar (mortar L) substrate. The plasters on brick substrate were tested either not painted or painted with silicone paint Funcosil.



Fig. 5.10 - Specimens (side view) used in set 1 tests: LNEC on brick (specimen 7LS), Parlumière on brick (specimen P8) and LNEC on lime mortar (specimen 11) without or with protective tissue

The plaster/substrate specimens (Fig. 5.10) have flat dimensions of 40 mm x 40 mm to 50 mm x 50 mm. MEP and Parlumière plasters are composed of a single mortar coat with a thickness of 20 mm. LNEC plaster is composed of a 5 mm adherence coat and a 20 mm base coat. Brick or lime mortar bases with a thickness of 20 mm or 30 mm, respectively,

were used as substrate. Preparation of the specimens is described in Annex II. At around the two months of age they were laterally sealed with an epoxy resin. The bottom face of the lime mortar substrate was protected with cotton tissue (Fig. 5.10) to avoid material loss during absorption or while handling the specimens.

NaCl or Na₂SO₄ crystallization tests were performed as reported in Table 5.3. During drying, the bottom faces of the specimens were sealed with polyethylene film.

		Salt solution and number of specimens used				
Materials	Paint	NaCl 1.9 molal (10%W)	Na ₂ SO ₄ 0.8 molal (10%W)	Pure water		
MEP-SP® on brick	Unpainted	3	2	3		
	Silicone paint	3	2	3		
Parlumière® on brick	Unpainted	3	2	3		
	Silicone paint	3	-	3		
LNEC on brick	Unpainted	2	2	2		
	Silicone paint	2	2	2		
LNEC on lime mortar	Unpainted	3	3	2		

Table 5.3 - Set 1 crystallization tests

The crystallization tests were carried out according to the following procedure (Vergès-Belmin *et al.* 2005):

- 1) The specimens are dried at 60°C in a ventilated oven and then allowed to cool in a conditioned room at around 20°C and 50% RH.
- 2) Wet-dry cycles are performed as follows:
 - 2.1) The specimens are partially immersed in the solution for 5 minutes, the liquid surface being around 5 mm above their bottom face. The 5 minute period was chosen so that:
 - the wet front nearly reached the plaster/brick interface;
 - the lime-mortar substrate absorbed a similar amount of solution.
 - 2.2) The specimens are subjected to daily cycles of 16 hours at 40°C and 20% RH plus 8 hours at 20°C and 50% RH in a climatic chamber with a low air velocity (the same FITOCLIMA 500 EDTU® chamber by Aralab used in the experiments of Chapters 3 and 4).
 - 2.3) The specimens are weighed, using a weighing device accurate to 0.001g, and the damage evaluated every week.
 - 2.4) The wet/dry cycle ends when at least 80% of the water that the specimens contained immediately after being contaminated has evaporated.

To prevent distortion of the results and since the drying behaviour of the two or three specimens in each group was always very coherent, it was decided to end each cycle only when all the specimens in the group had met this criterion. Indeed, since the specimens were weighed only once a week, minimal dephasement in their drying process would be magnified if the wet/dry cycle ended in different weeks. Differently, the last cycle was prolonged until the mass of the specimens was reasonably constant in time. There were two main reasons for this. Firstly, to minimize the amount of moisture present in the materials when they were sliced, not only because wet materials have a lower mechanical resistance but also because slicing induces always a certain increase in temperature which could cause migration of the liquid in the porous networks. Secondly, because, due to operational limitations, some specimens could not be immediately sliced and constant mass was a condition which allowed more flexibility as to the moment when the specimens are removed from the climatic chamber.

- 3) Each specimen is typically subjected to five wet/dry cycles, though, as will be seen, more than five cycles had to be carried out in some cases.
- 4) The salt distribution across the specimens is measured, at the end of the test, as follows:
 - 4.1) the specimens are sliced in a sawing machine at indents of typically 5 mm (the loss of material corresponds to a thickness of around 2 to 3 mm per cut);
 - 4.2) the cut samples are then crushed into powder in an agate mortar;
 - 4.3) the upper coat of the traditional plaster, however, could not be sawed due to its low cohesion and relatively large grain size; this mortar was split by progressive disagregation on sandpaper;
 - 4.4) the salt content in each sample is measured by means of the HMC method (at 20°C and 95% RH) using shallow-wide Petri dishes which, in this case, provided a sufficient accuracy;
 - 4.5) salt distribution profiles are obtained typically for one specimen of each type and each testing condition; in these profiles the HMC is plotted against the distance to the top surface of the specimen; each point in a profile represents the average HMC in that particular slice of the specimen.

5.5.3 - Results

5.5.3.1 - Salt crystallization tests

The results of the crystallization tests are presented in Tables 5.4 to 5.7. The behaviour of the two or three specimens in each group was quite similar as regards the amounts of salt introduced, drying rates and damage patterns. Hence, unless otherwise specified:

- damage description refers to all the specimens in the group;
- the damage patterns, normally illustrated by pictures of one specimen, are representative of the entire group of specimens;
- the salt distribution profile, which was typically measured for one specimen per group, is assumed to represent the salt-accumulation behaviour of the entire group;
- the total amount of salt reported in each distribution profile refers to that particular specimen but is similar for all the specimens in the group, as seen in Table 5.8.

Among these very coherent behaviours, LNEC plaster on lime mortar was an exception when tested with Na_2SO_4 (Table 5.7). Despite the amount of salt absorbed and the drying rate were, as further shown in Table 5.8 and Fig. 5.16, similar for the three specimens, only specimen 14 showed damage. The two remaining specimens (15 and 16) were subjected to additional wet-dry cycles but, even so, no visible damage occurred in neither of them.



Table 5.4 – Crystallization test on MEP-SP / brick specimens







Table 5.6 –	Crystallization	test on LNEC /	brick specimens
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Table 5.7 - Crystallization test on LNEC / lime-mortar specimens

5.5.3.2 - Tests with pure water

No damage was achieved in any of the tests with pure water. The distribution profiles of one specimen per group are presented in Fig. 5.11 and show insignificant HMC values.

As seen, some of the specimens were subjected to more than five cycles with pure water. The reason is that the tests with pure water served also to evaluate the stability of the environmental conditions and, hence, additional cycles had to be performed in some cases in order to accompany the tests with salt solutions which were, in general, much slower.



Fig. 5.11 - HMC profiles of some unpainted specimens tested with pure water

5.5.3.3 - Salt introduced in the specimens

Table 5.8 shows that quite similar amounts of salt were introduced in the tested specimens, particularly within each group.

		Salt introduced after 5 wet-dry cycles (g/m ²)							
	MEP-SP®	on brick	Parlumière® on brick		LNEC on brick		LNEC on lime mortar		
	Unpainted	Silicone paint	Unpainted	Silicone paint	Unpainted	Silicone paint	Unpainted		
NaCl	2416 2309 2356	2305 2256 2390	2373 2264 2275	2400 2295 2536	2249 2607	2373 2386	2182 2057 2112		
Na ₂ SO ₄	2020 2343	2091 2399	2151 2069	-	2351 1954	1989 2052	2236 2086 2264		

Table 5.8 - Salt introduced in the specimens

5.5.3.4 - Drying curves

The gravimetric drying curves obtained from weekly weighing the specimens are presented in the figures bellow. Figs. 5.12 to 5.15 concern the average values achieved with the two or three specimens of each type of material under each testing condition, which were similar to the individual values in every group of specimens.

Fig. 5.16 shows the individual drying curves of the three LNEC / lime-mortar specimens tested with Na_2SO_4 . As shown in Table 5.7, this was the only group of specimens where damage did not occur simultaneously in all the specimens. Fig. 5.16 shows that, nevertheless, the drying behaviour of the three specimens is very similar.

The tests with pure water served also, as mentioned, to evaluate the stability of the environmental conditions during the crystallization tests. That was done by checking whether the drying rate of the water-filled specimens changed from cycle to cycle. As seen in Figs. 5.12 to 5.15 this did not happen which indicates a good stability of the environmental conditions. Some minor changes observed between the first and the second cycles could be due to hydration reactions occurring at the first contact of the hardened mortars with free water.



Fig. 5.12 – MEP-on-brick drying curves



Fig. 5.13 - Parlumière-on-brick drying curves



Fig. 5.14 - LNEC-on-brick drying curves



Fig. 5.15 - LNEC-on-lime-mortar drying curves



Fig. 5.16 - Drying curves of the three LNEC-on-lime-mortar specimens with sodium sulfate

5.5.3.5 - Summary of results

The results of the tests carried out with NaCl, Na_2SO_4 or pure water are summarized in Tables 5.9 to 5.11, respectively.

				Durability	
Materials	Paint	Salt-accumulation behaviour	Alteration patterns	First observation of damage (weeks)	Test duration (weeks)
MEP-SP®	Unpainted	Accumulating (accumulation at the plaster mid-thickness)	Transverse cracks in the plaster 1cm from the surface	11	24
on brick Silicone	Silicone paint	Transporting	Efflorescence on all specimens; scaling of the paint layer on one specimen	6	24
Parlumière®	Unpainted	Accumulating (accumulation close to the interface with the substrate)	Transverse cracks (preceded by efflorescence) in the plaster at 2 mm from the interface	6 (efflor.) 11 (cracks)	17
on brick	Silicone paint	Accumulating (accumulation close to the interface with the substrate)	Transverse cracks (preceded by efflorescence) in the plaster at 2 mm from the interface	4 (efflor.) 13 (cracks)	22
LNEC on	Unpainted	Transporting	Efflorescence, sanding and peripheral cracks on the surface	4 (efflor.) 16 (cracks) 30 (sanding)	30
DIICK	Silicone paint	Transporting	Efflorescence with rupture of paint	4	33
LNEC on lime mortar	Unpainted	Accumulating (accumulation in the plaster mid-thickness)	Peripheral and diagonal cracks on the surface	14	22

Table 5.9 - NaCl cr	vetallization tests	(set 1)). enmar	v of regulte
	ystamzation tosts		j. Summar	y or results

Table 5.10 - Na₂SO₄ crystallization tests (set 1): summary of results

			Durability		
Materials	Paint	Salt-accumulation behaviour	Alteration patterns	First observation of damage (weeks)	Test duration (weeks)
MEP-SP®	Unpainted	Accumulating (accumulation in the half plaster next to the interface)	No alteration	-	20
on brick	Silicone paint	Accumulating (accumulation in the half plaster next to the interface)	No alteration	-	19
Parlumière® on brick	Unpainted	Accumulating (accumulation close to the interface)	Transverse cracks in the plaster 1 mm from the interface	11 to 13	19
LNEC on	Unpainted	Transporting (slight accumulation in the plaster)	Efflorescence	19	30
brick	Silicone paint	Transporting (slight accumulation in the plaster)	Efflorescence	35	35
LNEC on lime mortar	Unpainted	Accumulating (accumulation in the plaster mid-thickness + slight accumulation up to the interface with the substrate)	One specimen: longitudinal cracks in the plaster and transverse cracks at plaster/substrate interface	15	17

Materials	Paint	Test duration for 5 cycles (weeks)
MEP-SP® on brick	Unpainted	9
MEF-SF® ON BICK	Silicone paint	14 *
Parlumière® on brick	Unpainted	9
	Silicone paint	13
I NEC on brick	Unpainted	5
	Silicone paint	9
LNEC on lime mortar	Unpainted	7

	Table 5.11 -	Tests with	pure water (set 1): summar	v of results
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* only 4 cycles were performed; the 14 weeks were estimated for a 5-cycle testing period due to the constant cycle duration obtained in these 4 cycles (2, 3, 3 and 3 weeks)

5.5.4 - Discussion on set 1

5.5.4.1 - Main behaviour features

The behaviour of plasters and renders in relation to soluble salt crystallization can be described according to three main features: salt-accumulation behaviour, damage patterns and durability.

Salt-accumulation behaviour

The salt-accumulation behaviour of plasters and renders (Fig. 2.10) is perhaps the fundamental discriminating feature. It determines the depth(s) at which salts precipitate and, thus, the observance of major use requirements related, for instance, to the occurrence of surface damage or salt deposition at the plaster/substrate interface.

In the present crystallization tests, all plasters behaved either as salt-accumulating or as salt-transporting. Indeed, the salt-accumulation profiles in Tables 5.4 to 5.7 show that the salt crystallizes always either in or on the plaster. Nonetheless, only Parlumière presented constant salt-accumulation behaviour under different conditions:

- MEP and Parlumière both tended to behave like salt-accumulating plasters. Most salt was concentrated either at the middle-thickness of the plaster (MEP) or close to the plaster/substrate interface (Parlumière). The different penetration depth is probably due to the fact that the two plasters have similar vapour transport properties (Fig. 5.8) but Parlumière has lower capillarity suction (Fig. 5.5). Hence, the drying front, whose depth depends on a balance between the liquid and vapour transport, tends to be more deeply located in Parlumière.
- However, with NaCl, the painted MEP worked as a salt-transporting plaster.
- LNEC worked as salt-transporting or salt-accumulating plaster when applied on brick or on lime mortar substrate, respectively.

Damage patterns

The damage patterns are also relevant. As stated by Schaffer (1932), the type of decay is sometimes of even more significance than the decay rate. However, the damage patterns do not have to be considered always because they are closely related to the accumulation behaviour. Nonetheless, in some cases, they may allow a more detailed analysis of the performance of plasters and renders. Indeed, although typical damage patterns exist for each type of salt-accumulation behaviour, a given accumulation behaviour may give rise to

different damage patterns. For example, salt-transporting plasters tend to develop surface alterations (while salt-blocking plasters, for instance, are more likely to detach from their substrate). But these surface alterations may be varied in nature: efflorescence may or may not be accompanied by scaling, exfoliation, sanding or other.

The present crystallization experiments gave rise to different damage patterns, namely, surface damage or cracking. There are two main types of cracks: (i) transversal cracks are perpendicular to the central axis of the specimen; when they occur in the plaster, they correspond to detachment of part or of the whole plaster layer; (ii) longitudinal cracks are perpendicular to the plan of the plaster, that is, parallel to the central axis of the specimen; they are usually visible on the surface of the plaster as peripheral or diagonal cracks.

Transversal cracking or surface damage (efflorescence, sanding of plaster or rupture of the paint layer) are recurrently associated to salt-accumulating or salt-transporting behaviour, respectively. Differently, longitudinal cracking seems to be associated with a specific type of plaster (LNEC) rather than a certain salt-accumulation behaviour.

Durability

Durability is also a major feature, namely because of its economic significance. In fact, even if the salt-accumulation behaviour and damage patterns are similar, damage may occur at distinct rates. But, as stated by Knöfel *et al.* (1987), the test cycles in accelerated ageing tests can not be correlated to a period of field exposure, hence, straightforward prediction of the durability of building materials is not possible by means of these tests.

In the present experiments, the total amount of salt introduced after five wet/dry cycles is quite similar in all cases (Table 5.8) and, therefore, it is not a distortion factor when the durability of the various plasters is compared. However, comparing the durability of plasters with different accumulation behaviour is not a straightforward purpose. It is not feasible to base such a comparison on the intensity of damage because different damage patterns may occur and, hence, it would be necessary to rank them in a, certainly controversial, order of importance. It seems more reasonable to access the durability by considering the moment when damage is first observed, the intensity of damage being regarded as a second order factor. Still, two questions arise:

- Should surface alterations be considered as damage for salt-transporting plasters? To answer this question, efflorescence has to be distinguished from material damage. Indeed, it seems reasonable to disregard efflorescence as damage because transporting plasters are expected to carry salt solutions to the surface and, moreover, efflorescence can be removed. Yet, surface damage such as paint rupture or sanding has to be considered because it represents irreversible material damage.
- What if a certain plaster, which was aimed at working as, for instance, salt-accumulating, behaves as salt-transporting and, thus, shows efflorescence? In that case, it is logical that efflorescence is regarded as damage. In fact, salt-accumulating plasters are used mostly to avoid surface damage, for example, in houses. Hence, efflorescence corresponds, in this case, to a corruption of the fundamental working principle of the plaster.

Following this trend of thought, efflorescence was considered as damage for MEP and Parlumière but not for LNEC plaster. In any case, a comparison of the absolute durability of different plasters is hardly possible in the present experiments. Indeed, the durability of the plasters changed significantly for different boundary conditions (salt, paint or substrate material) as further discussed in sections 5.5.4.2 to 5.5.4.4.

Total duration of the test

It is also interesting to compare the total duration of the test, that is, the period necessary to achieve five complete wet/dry cycles (Tables 5.9 to 5.11).

Drying with pure water is, as expected, always faster than with the salt solutions. However, different rankings arise for pure water and the salt solutions as regards the drying speed of the different plaster/substrate systems. In fact, with pure water, the faster drying rate occurs for transporting plaster LNEC on brick. In contrast, with the salt solutions, the test is shorter for the plasters which tended to behave as salt-accumulating: MEP, Parlumière and LNEC on lime mortar. This suggests that, although pure water may be used as (a rough) reference for a certain material, it is not possible to compare the behaviour of different materials in relation to salt crystallization by reference to pure water.

5.5.4.2 - Influence of the type of salt

The total amount of salt introduced after five contaminations is quite similar in all cases, regardless of the type of salt (Table 5.8). This is in agreement with Hall and Hoff's (2002) statement that, in contrast to vapour transport, soluble salts do not significantly change liquid transport in porous materials.

However, sodium sulfate crystallizes always at greater (or, at least, similar) depths than NaCl (Tables 5.4 to 5.7). Even for the unpainted Parlumière, where both salts accumulated so close to the plaster/substrate interface that no depth differences are noticed in the salt-distribution profile, the transverse crack is closer to the substrate with Na₂SO₄ than NaCl (Table 5.5). The lower vapour pressure of NaCl is a possible cause since, as discussed in Chapter 4, it induces a more advanced position of the drying front. This difference can probably account also for the change in salt-accumulation behaviour of painted MEP: it worked as a transporting plaster with NaCl and as a salt-accumulating plaster with Na₂SO₄.

On the other hand, sodium chloride originates more localized (or, at least, similarly concentrated) accumulation of salt than sodium sulfate. The detail provided by the slicing technique is perhaps not great enough to allow, in some cases, for the observation of small distribution differences. Indeed, no relevant differences are observed between the NaCl and Na₂SO₄ distribution profiles of Parlumière (Table 5.5) or LNEC on lime mortar (Table 5.7). For painted MEP (Table 5.4) a comparison is also not possible because the accumulation behaviour was completely different with each salt. Nonetheless, a more localized deposition of NaCl than of Na₂SO₄ visibly occurred for unpainted MEP (Table 5.4) and also, although less clearly, for painted and unpainted LNEC on brick (Table 5.6). A more localized accumulation of NaCl by comparison to Na₂SO₄ could explain why the maximum salt content is, as seen in the distribution profiles (Tables 5.4 to 5.7), systematically higher for sodium chloride (HMC=350%~30% NaCl by weight) than for sodium sulfate (HMC=60%~13% Na₂SO₄ by weight).

It is not known whether the temperature cycles contribute for the more even distribution of sodium sulfate. Indeed, the drying cycles have a 40°C period (16 hours) and a 20°C period (8 hours). Above 32.4°C, thenardite is the only equilibrium phase of sodium sulfate. Therefore, thenardite is the equilibrium phase at the end of each 40°C period, when a new 20°C period starts. However, in contact with a sodium sulfate solution below 32.4°C, mirabilite is always the equilibrium phase (McMahon *et al.* 1992, for instance). This happens in the wet zones of

the specimens during the 20°C drying periods. Therefore at each transition from 40°C to 20°C, the equilibrium phase in solution varies from thenardite to mirabilite. Yet, the solubility of thenardite is much higher than the solubility of mirabilite (Fig. 2.8). Therefore, temperature-induced crystallization of mirabilite may readily follow each 40°C-20°C transition, regardless of the position in the specimen, hence, not necessarily at the drying front.

Crystallization of mirabilite may also arise at the contact of the salt solution with pre-existing crystals. Indeed, the stable equilibrium phase in the air at either of the drying conditions (40°C and 20% RH or 20°C and 50% RH) is thenardite. Therefore, an increasingly significant amount of crystalline thenardite is expected to be present in the pore system of the plaster during each drying period, as the specimen becomes dryer. At the contact with the liquid solution introduced by a new contamination, thenardite dissolves giving rise to a solution which is highly supersaturated with respect to mirabilite. Hence, mirabilite can crystallize readily. This sort of process was reported by Chatterji and Jensen (1989) and also, more recently, by others (Tsui *et al.* 2003, for instance).

Unlike usual in crystallization tests, NaCl caused more severe damage than Na_2SO_4 in the present experiments. This happened probably because NaCl tends to crystallize in a narrower volume. Although higher crystallization pressures are often attributed to Na_2SO_4 (Rodriguez-Navarro and Doehne 1999 for instance), a higher pressure per unit volume may thereby arise for NaCl. Indeed, as seen in Tables 5.4 to 5.7:

- In the case of salt-transporting behaviour, NaCl efflorescence is always accompanied by material damage: rupture of paint layer for MEP and LNEC on brick; longitudinal (peripheral) cracking and sanding of the plaster for unpainted LNEC on brick. In contrast, Na₂SO₄ induced no surface damage other than efflorescence.
- In the case of salt-accumulating behaviour, cracking arises systematically with NaCI: transverse cracks at the depth of major salt-accumulation occurred in MEP and Parlumière; longitudinal (peripheral and diagonal) cracks arose on LNEC/lime-mortar. Yet, with Na₂SO₄ damage was observed only in some of the salt-accumulating plasters: Parlumière and one out of the three LNEC/lime-mortar specimens.
- Longitudinal (peripheral or diagonal) cracking occurred only for unpainted LNEC. But also as regards this kind of damage NaCl was more harmful than Na₂SO₄. With NaCl, all the specimens on both substrates cracked, that is, cracks occurred either in the case of salt-transporting or salt-accumulating behaviour. With Na₂SO₄ only one of the LNEC/lime-mortar specimens developed longitudinal cracks. None of the remaining unpainted LNEC specimens, either on lime mortar or on brick substrate, cracked with Na₂SO₄.

Another aspect of the higher damage capability that NaCl revealed in this set of tests is that the overall durability of the plasters is lower or, at the best, similar with NaCl than with Na₂SO₄. As seen in Tables 5.9 and 5.10, damage occurs always first with NaCl.

Yet, the difference between the two salts is not very significant when duration of the tests is considered. A faster drying process was expected for Na_2SO_4 by comparison to NaCl because the RH_{eq}^{sat} of Na_2SO_4 is higher than that of NaCl (Table 5.12). Yet, although in some cases (MEP and LNEC on lime mortar) a somewhat shorter testing period indeed arises for Na_2SO_4 , in other cases (Parlumière and LNEC on brick) this period is similar for both salts or slightly shorter for NaCl.

Although salt deposition and cracking of the specimens can certainly influence the drying rate, this unexpected drying behaviour is likely to arise also from the fact that, in these tests, only the substrate was saturated. Hence, a slightly deeper location of the drying front for Na_2SO_4 could, to some extent, compensate the influence of a higher vapour pressure on vapour transport during drying.

Solution		Relative equilibrium humidity RH _{eq} (%)			
		At 20°C	At 40°C		
NaCl	10%W (1.9 molal)	94.3 ⁽¹⁾	94.2 ⁽⁶⁾		
	Saturated	75.5 ⁽²⁾ (6.1 m)	74.7 ⁽²⁾ (6.1 m)		
Na ₂ SO ₄	10%W (0.8 molal)	97.5 ⁽¹⁾	97.4 ⁽⁶⁾		
	Saturated	95.6 ⁽³⁾⁽⁵⁾ (3.5 m)	≈ 86 ⁽⁴⁾ (3.3 m)		

Table 5 12 Dale	tivo oquilibrium bumidit	v of aquaqua	NoCl or No. SO	colutions at 20	10C or 400C
1 able 0.12 - 1 cla	live equilibrium numun	ly of aqueous i	11401 01 1142004	Solutions at Z	J C 01 40 C

(1) at 25°C, from Robinson and Stokes (2002)

(4) from Tsui *et al.* (2003); refers to thenardite(5) refers to mirabilite(6) from Reis (2006)

Indeed, as seen in the drying curves of Figs. 5.12 to 5.15, during the first cycles the drying rate is in all cases very similar for pure water and for both salt solutions. The further decrease in the drying rate probably arises more from the deposition of salt than from vapour pressure differences. This agrees to the conclusion in Chapter 4 that the deposited salt crystals play a major part in hindering drying. It is interesting that the drying rate of LNEC plaster (on both substrates) decreases up to the 3^{rd} cycle and then stabilizes or rises again. In some cases this change clearly coincides with visible cracking of the specimens (unpainted LNEC on brick and LNEC on lime mortar with NaCl). This does not happen for LNEC on lime mortar with Na₂SO₄ where cracking is observed only in one specimen (Table 5.7) and, yet, the drying rate is similar for the three specimens. Therefore, the reasons for the further rise or stabilization of the drying rate are not totally clear. One possibility is the occurrence of micro-cracks in the two other LNEC/lime-mortar specimens, which were not detected by visual observation.

5.5.4.3 - Influence of the paint

Tables 5.4 to 5.7 and Figs. 5.12 to 5.14 show that the drying rate in the first cycles, before cracking was observed, is in general slightly lower for the painted specimens, both in the case of pure water and the salt solutions.

It is interesting that for MEP and Parlumière, which have similar vapour permeability (Fig. 5.8), the use of paint induces a similar increase in the duration of the tests with pure water from 9 to 13 or 14 weeks (Table 5.11). LNEC's vapour permeability is higher but the drying rate decreased less in the presence of the paint (5 to 9 weeks). This could suggest that the conclusion of Hern and Snethlage (1992) that the more vapour permeable a stone, the higher the resistance to vapour transport induced by certain paint, was not verified here.

⁽²⁾ from Greenspan (1977)

⁽³⁾ from Linnow et al. (2006)

However, such direct comparison can probably not be made because LNEC dries fundamentally by surface evaporation while for Parlumière and MEP the drying front is located inside the plaster.

With NaCl, the paint changed the salt-accumulating MEP into a salt-transporting plaster. In contrast, no significant (Parlumière) or no visible (MEP with Na₂SO₄ and LNEC) alteration of the accumulation behaviour arose for the other plasters through the use of paint. If for MEP with Na₂SO₄ or LNEC it is not impossible that some minor differences were masked by an insufficient detail provided by the slicing method, for Parlumière it is clear that the transverse cracks occurred at similar depths with and without paint.

These differences are not easy to interpret without the help of a proper numerical model or of an efficient monitoring technique such as MRI. Drying, especially in the presence of salt, is a complex dynamic process where liquid transport also plays a role. Moreover, the cracks, which may only be detected at advanced stages, may hinder liquid moisture transport or enhance evaporation.

Nonetheless, a general conclusion is possible: a paint may in certain conditions transform a salt-accumulating plaster into a salt-transporting plaster, although this will not necessarily happen for all plasters or salts.

When damage arose for a certain plaster, the use of paint speeded up its occurrence. It happened for MEP, Parlumière and LNEC on brick with NaCl. The paint caused a poorer performance of the salt-accumulating plasters MEP (that changed into salt-transporting) and Parlumière (whose transverse cracks probably arose earlier since efflorescence on that area was first observed on the painted specimen). But salt-transporting plaster LNEC on brick may be a different case. Indeed, although NaCl damage arose earlier on the painted specimens, it was less severe (efflorescence and further rupture of the paint layer) than on the unpainted specimens (longitudinal cracking).

5.5.4.4 - Influence of the substrate material

It is clear that the substrate material can greatly affect the behaviour of a plaster in relation to salt crystallization. Indeed, despite the fact that five wet/dry cycles were performed and similar amounts of salt were introduced in each case, LNEC behaved as a salt-transporting plaster when applied on brick substrate (Table 5.6) and as a salt-accumulating plaster when applied on lime mortar substrate (Table 5.7).

As regards durability and damage patterns, LNEC seems to have an overall tendency for cracking, as discussed above. Nonetheless, cracking is more severe in the case of lime mortar substrate: for NaCl, cracking arose first and was more intense on lime mortar substrate; for Na₂SO₄, one of the specimens with lime mortar substrate cracked but none of those with brick substrate did.

The reasons for this different behaviour will be discussed in section 5.10.2 considering also results obtained in further sets of tests.

5.5.4.5 - Test method

There is no doubt that the present crystallization tests were shorter than those reported by van Hees and Brocken (2004). Here, a maximum duration of 17 to 35 weeks was achieved for the different plasters in contrast to the 12 to 48 months in that research. However, even 17 weeks is not a short period. It can be acceptable perhaps for long-term research studies but hardly when plasters are evaluated in support of consultancy or product development studies.

A second important question that arises on the test method concerns the definition of its end point. In the present experiments every specimen was subjected to five wet/dry cycles, which allowed carrying out of sustained comparisons between materials with similar amounts of salt. Yet, cracks arose in some specimens. Hence evaporation could have been enabled (Vergès-Belmin *et al.* 2005), namely lateral evaporation through transverse cracks that eventually broke the epoxy sealing, or liquid transport hindered by those cracks. The significance of these effects as regards test representativity is difficult to evaluate. Yet, although it is possible that cracks occur before being visible, it seems reasonable in future tests to slice one specimen of each kind at the first observation of damage. Furthermore, because damage represents a failure condition, it may be useful to know the associated salt distribution.

Another relevant concerns the slicing technique. In most cases, a sawing machine was used which originated a loss of material correspondent to a thickness of around 2 to 3 mm per cut. Although a discrete salt distribution can be perfectly acceptable, it must be kept in mind that a significant percentage of salt is lost by this technique. Therefore, quantitative comparisons should be made cautiously especially because some salts, such as NaCl, may deposit in a relatively narrow band. In extreme situations, which were not the case here as seen in the distribution profiles in Tables 5.4 to 5.7, most salt may even be lost. Therefore, whenever possible, sampling techniques that do not involve material loss should be used. Such a technique was used here on the traditional mortars. These mortars were divided by manual sandpapering which, however, is a technique that requires too much time and effort to be practicable in large groups of specimens (here, the 50 specimens tested in total originated almost 400 samples).

Finally, the HMC method by means of which the distribution profiles were obtained could have provided inaccurate results in the case of sodium sulfate, as it happened in some of the experiments described in Chapter 3. Indeed, the set-point RH was here also of 95%, hence, below 95.6%, the newly proposed value for the deliquescence RH of mirabilite at 20°C (Linnow *et al.* 2006) which the experiments presented in section 3.5.4.5 corroborate. Therefore, the HMC values obtained in the present set of tests were carefully considered. It was found that the distribution profiles were very coherent among the various sulfate-loaded specimens and between these and the chloride-loaded specimens, both in terms of distribution and maximum HMC values. It is therefore likely that the actual RH in the chamber was somewhat above 95.6% during these HMC measurements.

In this case no control-samples were used to monitor the environmental conditions in the chamber because the conclusions of the HMC experiments presented in Chapter 3 had not yet been achieved at the time these crystallization tests were performed.

5.6 Drying of a salt-accumulating plaster (set 2)

5.6.1 - Introduction

This section presents several MRI-monitored drying experiments performed on MEP/brick specimens loaded with NaCl or Na₂SO₄. Unpainted MEP on brick substrate was used because it behaved as an accumulating plaster in set 1 crystallization tests. Indeed, salt-accumulating plasters are of high practical interest. Hence, further insight into their behaviour will hopefully contribute to enabling the development of increasingly effective products.

Parlumière also worked as a salt-accumulating plaster but the salt accumulated very close to the plaster substrate interface, approaching somewhat the behaviour of a salt-blocking plaster. In contrast, in MEP, the salt penetrated up to the middle-thickness of the plaster. Moreover, very clear differences between the behaviour with NaCl and Na₂SO₄ were observed in unpainted MEP: NaCl accumulated in a narrow band and caused transverse cracking of the plaster; in contrast, Na₂SO₄ was more evenly distributed between the plaster/substrate interface and the middle-thickness of the plaster and caused no visible damage.

Drying experiments where only the substrate was contaminated, as well as experiments on saturated specimens, were carried out. The main objectives of this research were the following:

- understanding liquid migration features underlying the accumulation behaviour of MEP plaster;
- comparing the behaviour of this plaster when only the substrate is contaminated to that arising in the case of capillary saturation of the whole plaster/substrate specimen.

5.6.2 - Materials and methods

The drying experiments were performed on MEP/brick specimens filled with pure water, 1.9 molal (10%W) NaCl solution or 0.8 molal (10%W) Na₂SO₄ solution.

For pure water and NaCl two different contamination conditions were used: (i) capillary saturation of the whole specimen by total immersion during an overnight; (ii) contamination of the substrate only, by maintaining the bottom 5 mm of the brick immersed in the solution, for some minutes, until the wet fringe attained the brick/plaster interface. For Na_2SO_4 only contamination condition (ii) was used.

The 1D MRI technique used to monitor these experiments (Annex I) provides profiles that express the concentration, in arbitrary units (a.u.), of the H^+ ion over the length of the specimen (Fig. I.4). Each profile corresponds to a different moment of drying, hence, the different profiles obtained within one drying experiment represent the evolution over time of the moisture content across the specimen.

The experiments were carried out on cylindrical specimens (Fig. I.2) drilled from prismatic specimens similar to those of set 1 crystallization tests (Fig. 5.10). The specimens were bottom and laterally sealed during drying to ensure unidirectional migration of moisture. Drying proceeded under airflow at 0%RH (temperature \approx 18°C). In all the experiments: (i) a first profile was obtained before the airflow was turned on; (ii) the airflow was turned on 30 minutes before the second profile was obtained.

At the end of the experiments, the specimens were oven dried at 40°C. The salt distribution was then measured across specimens MEP 1 and MEP 2, the ones where only the substrate was filled with sodium sulfate or sodium chloride solution, respectively. The salt distribution profiles of MEP 2 have to be carefully considered because a second experiment, performed under similar conditions but using a more concentrated (4 molal) NaCl solution, was carried out on this specimen. This second experiment, which is not presented because it was very short and not conclusive, was performed for figuring out whether an acceptable definition of the Na⁺ ion could be achieved for MEP plaster (which did not occur).

In order to obtain the salt-distribution profiles, the specimens were divided at indents of typically 5 mm by means of progressive sandpapering. Afterwards, the salt content in each of the obtained samples was measured. First, HMC tests were performed in a climatic chamber set pointed at 20°C and 95% RH, using Petri-dishes. Coherent HMC profiles resulted for NaCl but the samples contaminated with Na₂SO₄ achieved only the HMC of each base material (around 5% or 3% for the mortar and the brick, respectively). Control samples of pure NaCl, tested simultaneously, achieved a HMC of around 740% which corresponds to an RH of 92% at 20°C. This indicates that the actual RH in the chamber was far below the RH_{eq}^{sat} of mirabilite (95.6%) which could not thus deliquesce. Yet, this actual RH was well above the RH_{eq}^{sat} of thenardite (87%) and no hydration took place either, which indicates a metastable behaviour of sodium sulfate in this case. The salt content in the same samples was then measured (Esteves and Catarino 2006) by ion chromatography (IC). IC was feasible in this case due to the reduced number of samples. It is interesting to see (Fig. 5.17) that a very good correlation HMC / ion content resulted for NaCl, despite the slightly different HMC of the two base materials involved.



Fig. 5.17 – Correlation HMC / ion content (the light and dark green points correspond to brick and mortar samples, respectively)

5.6.3 - Results

The results of the five drying experiments on saturated or partially saturated specimens are shown in Figs. 5.18 and 5.20, respectively. Each profile in these graphs is assigned to a time value that corresponds to the moment when the profile began being measured. The initial profile (red dashed line) is that obtained before the airflow was turned on. The 0h profile (red continuous line) is the first profile measured after the airflow being turned on. The first 20h of drying are always represented by profiles (black lines) at intervals of 2h. The subsequent drying period is illustrated with profiles (blue lines) at intervals of 10h. The last profile obtained in each experiment is depicted in pink. The top face of the plaster, through which drying took place, is located on the left part of the profiles.

The physical limits of the test specimens are not straightforward to identify in the NMR profiles. First, there is an uncertainty associated to the resolution at which the measurements were carried out (1 or 2 mm). Also, the bottom surface is not perfectly orthogonal, which may explain the progressive lowering on the measured values towards the edge of the three specimens in Fig. 5.20. Artifacts that consist in the further rise of the NMR signal values are observed near the bottom surface of the specimens in Fig. 5.20 b and c. They probably correspond to some water that was trapped in the Teflon sealing, which is consistent with the fact that the signal peak remained constant throughout the experiments. A similar effect could explain the signal peak visible in Fig. 5.18 a) at the 50 mm. Yet, in this case, the water was probably trapped next to the substrate material.

The top edges of the three partially saturated specimens in Fig. 5.20 are indistinct because the liquid penetrated only up to around the half thickness of the plaster and, hence, flat NMR profiles are obtained for the top part of the plaster. Nevertheless, the physical limits of these three specimens could be identified (they correspond to the grey vertical line segments) on the basis of a careful measurement of their dimensions. This was not possible for MEP5 specimen in Fig. 5.18 due to operational limitations (the last NMR measurements with NaCl solution were completed after the author's stay at TUE had finished). The possible location of the edges of this specimen is identified by the grey shadow areas in the figure.

The drying curves in Figs. 5.19 and 5.21 show the evolution over time of the total amount of liquid in the specimens. The total amount of water present at each moment corresponds to the area below the corresponding MRI profile. The drying rate is the slope of the drying curve.

Figs. 5.22 to 5.24 concern the liquid flux and evaporation flux to and from the drying front, respectively, during drying of the partially saturated specimens (Fig. 5.20). The liquid flux was assumed to correspond to the decrease in the moisture content of the brick substrate. Hence, it was calculated by considering the difference between the areas (within the brick) below two consecutive MRI profiles and, then, dividing it by the corresponding time interval. The evaporation flux was obtained in a similar way but considering the total area (within the whole specimen) below two consecutive profiles.

In the drying curves and flux graphs, time zero corresponds to the first profile obtained after the air flow was turned on. The profile obtained before the airflow was turned on corresponds, hence, to a negative time value.

In Figs. 5.25 and 5.26 the salt distribution profiles of the partially saturated specimens obtained by IC are presented. Additional IC measurements carried out on the non-contaminated materials indicate that MEP plaster contains a certain amount of sulfate. The sulfate content in the non-contaminated materials is represented by the dashed line in Fig. 5.26.



Fig. 5.18 - Results of the MRI-monitored drying experiments on fully saturated specimens



Fig. 5.19 - Drying curves: total amount of water in the saturated specimens



a) MEP6 - water (duration 20 hours)





c) MEP1 - Na₂SO₄ solution (duration 30 hours)

Fig. 5.20 - Results of the MRI-monitored drying experiments on partially saturated specimens



Fig. 5.21 - Drying curves: total amount of water in the partially saturated specimens



Fig. 5.22 - Difference between evaporation and liquid fluxes in the partially saturated specimens



Fig. 5.23 - Liquid fluxes in the partially saturated specimens







Fig. 5.25 - Chloride distribution in MEP 2



Fig. 5.26 - Sulfate distribution in MEP 1

5.6.4 - Discussion on set 2

5.6.4.1 - Totally saturated specimens

The results of the drying experiments on totally saturated specimens are shown in Fig. 5.18. For both pure water and the NaCl solution, MEP behaved as a transporting plaster, most of the water being eliminated by surface evaporation. The clear differences that arise between drying in the presence of pure water or of the salt solution are similar to those observed in the experiments of Chapter 4:

- Drying is much faster with pure water: approximately 70% of the water evaporates after 40h of drying while, for the salt solution, around 65% evaporates after 160h.
- Stage I is shorter for pure water. In this case, the surface starts almost immediately drying faster. The drying front recedes completely into the material after around 20h. In contrast, for NaCl, evaporation is still proceeding at the surface after 160h and no signs of a transition from Stage I to Stage II are visible yet, though the moisture content in the plaster is already far bellow that at which stage II begins for pure water.
- For pure water, a lower amount of moisture is eliminated by surface evaporation. In this case, approximately 60% of the water is eliminated at the surface, while for NaCl about 65% had already evaporated at the surface when, after 160 h, stage I is still apparently far from ending.

The moisture content decreases at a faster rate in the brick than in the plaster during almost the whole experiments. This indicates that the capillary suction of a significant percentage of the pores in the plaster is higher than the capillary suction of the pores in the brick (it is not reasonable to correlate capillary suction and pore size because the plaster includes hydrophobic additives). The drying rate of the brick becomes lower than that of the plaster in the last hours of the experiments. Yet, at the end of these experiments, approximately 70% (pure water) or 80% (NaCl) of the water initially present already left the brick. Hence, it is possible that, over a longer period, practically all the solution in the brick would have migrated to the plaster as liquid.

5.6.4.2 - Partially saturated specimens

The MRI profiles in Fig. 5.20 show that absorption and drying occur simultaneously during the drying experiments where only the substrate was contaminated:

a) Initially, the solution is mostly in the substrate. In all specimens, high and reasonably uniform moisture content exists in the brick at this moment. This initial moisture content is lower for the salt solutions than for pure water, largely because part of a salt solution is not water. Between the two salt solutions, the initial moisture content is lower for Na₂SO₄ than for NaCl, which is probably due to the contamination procedure being only roughly accurate. Material heterogeneity among the several specimens could also account, to some extent, for these differences. But the initial moisture content in the brick is lower in the partially saturated specimens than in the totally saturated specimen (Fig. 5.18). This suggests that not all the pores in the brick were filled when the wet front reached the plaster/substrate interface of the partially saturated specimens.

- b) Evaporation takes place on the wet front which, in the beginning, is located more or less at the plaster/brick interface as shown by the initial profiles (red dashed curves).
- c) Immediately after the brick is contaminated, the plaster starts sucking the liquid. Hence, the moisture content in the brick begins decreasing. It is interesting to note that the water content in the plaster is, next to the interface, approximately constant in the course of the experiments. This suggests that MEP pores involved in drawing water from the brick are always the same.
- d) The total moisture content in the specimens decreases as evaporation proceeds. However, the MRI technique does not seem to provide acceptable quantitative values for the early hours of the absorption/drying process. It can be seen, for instance in Fig. 5.21, that very low or even negative drying rates are measured at the beginning of the experiments. Yet, it is not likely that a very low evaporation rate exists precisely when the moisture content at the drying front is higher. Furthermore, negative rates would mean that the specimen was absorbing moisture which is not possible at 0% RH. In fact, those occurrences correspond to an experimental error. The key point is that the measurement of one profile takes 0.5 or 1.5 h (in the case of pure water or salt solutions, respectively). This is acceptable when only vapour fluxes are involved, as seen in the previous experiments on totally saturated specimens because the moisture content varies much more slowly. However, in the partially saturated specimens there is a simultaneous absorption process going on. A high amount of liquid is moving from the brick to the plaster while profiles are being measured by indents in the reverse direction. Hence, there is always a significant amount of moisture that is not detected: it has not reached the upper part of the specimen when this part is being measured and has already left the lower part when, later, measurements begin on this other part. Thus, total moisture contents lower than the actual ones result. The error seems to be higher for pure water than for NaCl and higher for NaCl than for Na_2SO_4 . This is logical because the higher the initial moisture content in the brick the higher the liquid flux, thus, the higher the error. This error affects also the flux graphs in Figs. 5.22 to 5.24.
- e) The wet front progresses in the plaster and eventually stabilizes several millimetres away from the outer surface, as seen in Fig. 5.20. Up to the end of the experiments no further movement of the front is observed, neither forwards or backwards.

At this point, a few questions arise. Why does the wet front stabilize? Why does it not recede with decrease in the total moisture content in the specimen? For example, in the case of NaCl which was the longer experiment, Fig. 5.20b shows that practically no movement arises from the 15h to the 60h (when the test ended) despite 30% of the total amount of water initially present having evaporated during that period.

What happens is probably the following. At the beginning of the drying experiments, the wet front is coincident with the plaster-substrate interface. There is a very steep gradient of the capillary potential because the plaster is dry immediately ahead of the interface. As indicated by Eq. 5.2, a very high liquid flux u arises and, so, the wet front moves rapidly into the plaster.

The wet front is not a sharp front but a diffuse front of width *e*, as seen in Fig. 5.20 and represented in Fig. 5.27. This happens probably due to the large range of pore sizes in MEP plaster (Fig. 5.7) and also to the presence of hydrophobic additives. It is likely that, for these

reasons, MEP plaster possesses several families of pores through which water migrates at different velocities. The capillary absorption tests are consistent with this hypothesis: in contrast to a common material such as brick, the absorption curve of MEP (Fig. 5.5) includes several line segments with distinct inclinations that indicate distinct water migration kinetics (Mertz 1991).



Fig. 5.27 – Representation of liquid migration in MEP plaster

Evaporation occurs in the diffuse wet front. It is interesting that, probably because of the combined effects of evaporation and differential liquid migration in the pores of MEP plaster, there is not exactly an advancement of the wet front, but rather an extension. It is also interesting that the moisture content in the plaster varies almost linearly across the diffuse front (Fig. 5.20).

The liquid flux decreases (Fig. 5.23), as the width of the wet front increases, due to lowering of the capillary potential gradient (Eq. 5.1). The vapour flux tends to increase as the wet front approaches the surface and, oppositely, to decrease with the lower availability of liquid at the wet front. It is not known how much the deposited salt crystals contribute to lowering the evaporation rate. The result of these opposed tendencies is probably an overall reduction of the vapour flux, as roughly suggested by Fig. 5.24. Yet, reduction of the vapour flux is lower than that of the liquid flux (Fig. 5.22). Eventually, the two fluxes meet in value and, so, the wet front stabilizes (Fig. 5.22). Up to the end of the three experiments, no further movement of the front is observed.

This steady state would be broken only by a decrease in the capillary potential next to the interface. This could occur if the moisture content in the brick decreased to a point where some of the pores in the plaster were no longer able to draw water from the brick. In the case of both pure water and Na_2SO_4 the test was too short for that. However, for NaCl, a very small decrease in the moisture content in the plaster is observed at the interface from the 18h onwards (Fig. 5.20b). And, yet, not even a slight receding of the drying front was observed.

This happens probably because advancement of the drying front corresponds to wetting of the plaster, while receding of that front corresponds to drying. Due to the hysteretic nature of absorption and drying processes (Fig. 5.4), at a given moisture content, the capillary potential is higher for drying than for wetting. Hence, the moisture content at the interface had to decrease further to account for that difference. Backwards movement of the drying front in this kind of processes is probably also limited by reduction of the vapour flux, due both to the further increase in the path length to the surface and to a decrease in the liquid flux.

None of the specimens in the present experiments ever attained the phase at which the drying front recedes: constant liquid (Fig. 5.18) and vapour (Fig. 5.24) fluxes were measured during the last period up to the end of the experiment.

Nevertheless, the considerations above suggest that a strong absorption/drying hysteresis can account for a longer stability of the drying front, thus, for a lower risk of salt being deposited on or in the substrate as a result of diminishment of the liquid flux.

It is interesting that stabilization of the drying front occurred at very similar depths in all the partially saturated specimens, as suggested by the profiles in Fig. 5.20. This is consistent with the low influence of soluble salts, at least up to the present concentrations, on liquid transport (Hall and Hoff 2002). Indeed the liquid flux determines the progression of the front during the early moments, as discussed above.

The drying rate (Fig. 5.21) is somewhat lower for NaCl than for Na_2SO_4 and for Na_2SO_4 than for pure water. These differences are consistent with the slightly different RH_{eq} of the three liquids, when either the original (10%W) or saturated salt solutions are considered (Table 5.12).

What was not fully expected in these drying rates is that they were practically constant over long periods in the experiments with salt solutions. Indeed, when the drying font is stable, if both the increase in the concentration of salt solutions and salt deposition lower the evaporation rate, the drying rate is expected to progressively decrease. Hence, there is not a clear explanation for these constant drying rates which shows the need for further research on the influence of salt crystals on porous material drying.

The final salt distribution profiles are shown in Figs. 5.25 and 5.26. These profiles have to be carefully considered because several possible error factors exist:

- Firstly, the specimens still contained a significant amount of solution at the end of the drying experiments. Therefore, part of the salt crystallized during the subsequent oven drying at 60°C. Such very fast drying is expected to cause a higher amount of salt to be dispersed all over the pore network, as discussed in Chapter 4. This is compatible with the fact that salt was deposited in the brick, despite the wet front being always in the plaster during the drying experiments. Indeed, in both specimens, the salt was quite evenly distributed in the brick, which agrees with the homogenous moisture distribution that existed in this material at the end of the drying experiments.
- Secondly, the specimens were divided only after several months. Therefore, further migration of salt might have arisen if some residual amount of moisture remained in the specimens or was attracted by hygroscopicity. Liquid transport towards the outer surface or diffusive (liquid or ion) homogenization processes might have occurred in this case. That could explain why some salt was found in the surface layers of the two specimens despite, in the drying experiments, the wet front never having reached those advanced positions.

- Thirdly, MEP 2 specimen was subjected, as explained in section 5.6.2, to a second contamination with a more concentrated NaCl solution. The higher salt load accounts therefore for the much higher ion content in the NaCl loaded specimen (Fig. 5.25).
- Finally, the sulfate content in MEP plaster may have influenced the salt distribution in Fig 5.26.

Despite these interferences, clear differences arose between the distributions of the two salts: NaCl deposited predominantly at the middle-thickness of the plaster while Na_2SO_4 precipitated mostly next to the interface with the substrate. These distinct deposition patterns, which are similar to those achieved in set 1 crystallization tests, can be due to solubility differences between NaCl and Na_2SO_4 :

- At 20°C, sodium chloride has only one crystal phase (halite), which readily crystallizes from solution at 6 molal (Fig. 2.6). At the same temperature, sodium sulfate can precipitate from solution in two different crystal phases, the decahydrate mirabilite or the metastable heptahydrate, whose maximum solubility is 1.7 molal and 3.15 molal, respectively (Fig. 2.8).
- At 40°C, the solubility of the equilibrium crystal phase of sodium sulfate (thenardite) is of around 3.5 molal, hence, lower than the solubility of halite which is approximately 6 molal also at this temperature.

Because evaporation occurs all over the (diffuse) drying front in the partially saturated specimens, the solutions become more concentrated as they migrate across the width *e* of this front (Fig. 5.27). Since, at 20°C, the solubility of sodium sulfate is lower than that of sodium chloride (Figs. 2.6 and 2.8), sodium sulfate can precipitate earlier in the drying front, hence, deeper in the specimen, as it happened in the present experiments (Figs. 5.25 and 5.26).

It is likely that also in set 1 crystallization tests a diffuse rather than a sharp front arouse during drying for MEP and possibly also for Parlumière and LNEC plasters. Hence, apart from the hypothesis put forward in section 5.5.4.2 that concern crystallization due to temperature changes or to contact between the salt solution and thenardite crystals, solubility differences among NaCl and Na_2SO_4 can also account for the more even and deeper deposition of the last:

- Sodium sulfate can, as stated, precipitate earlier in the wet front because the solubility of any of its possible crystal phases is significantly lower than that of halite both at 20°C and at 40°C.
- Sodium sulfate can crystallize from a wider range of concentrations than sodium chloride because it has two possible crystal phases and can easily supersaturate. Therefore, it may deposit at different points across the drying front. In contrast, sodium chloride has only one crystal phase and a low tendency to supersaturate. Hence, it will deposit in a more limited area.

5.7 Influence of a paint on drying (set 3)

5.7.1 - Introduction

MRI-monitored drying experiments were performed on painted or unpainted stone specimens. The main objective of those experiments was to evaluate the influence that paints may have on drying and, thus, on evaporative salt deposition processes.

5.7.2 - Materials and methods

The drying experiments were performed on two stone D specimens, one of which was painted with a commercial acrylic emulsion paint (Table 5.1). The experiments were monitored by means of the 2D MRI technique described in Annex I.

The specimens had flat dimensions of 20 mm x 30 mm and a thickness of 20 mm. The acrylic paint was composed of two coats which were applied with a 24 h interval.

Immediately before the experiments the specimens were lead to capillary saturation by total immersion in pure water for one hour. Afterwards, their lateral and bottom faces were sealed with Parafilm® plastic film. The specimens were then assembled together with Teflon tape (Fig. 5.28). This allowed the drying experiments to be performed simultaneously in both specimens.

Drying was carried out under airflow at 0%RH (temperature \approx 18°C). In all the experiments: (i) an initial image was obtained before the airflow was turned on; (ii) the airflow was turned on immediately before the second image started being measured.



Fig. 5.28 - Representation of the specimen assemblage (side-view) and cross-sections of interest

5.7.3 - Results

The results of the drying experiments are presented in the four figures below. They include a selected series of MRI snapshots (Fig. 5.29), 1D profiles (Fig. 5.30) at the two cross-sections indicated in Fig. 5.28 and the overall drying curves (Fig. 5.31). Each point on a drying curve corresponds to the sum of all values obtained across the specimen in the respective 1D profile.



Fig. 5.29 - MRI snapshots of specimens D2 and D1P during drying with pure water



Fig. 5.30 - One-dimensional profiles at crosssections NP (above) and P (below) indicated in Fig. 5.28. The grey vertical lines represent the physical limits of the specimens (the drying surface corresponds to the segment on the left). The green lines indicate the maximum penetration of the drying front in the material.



Fig. 5.31 - Drying curves. The shadow areas indicate the instants when, as seen in Fig. 5.30, the drying front could have receded totally into the material.

5.7.4 - Discussion on set 3

Both specimens go through a multistage drying process. During Stage I, the drying front is located at the outer surface (Fig. 5.29), the distribution of moisture is reasonably homogenous throughout the specimens (Fig. 5.30) and the drying rate is mostly constant (Fig. 5.31). Transition to Stage II begins when the moisture content starts decreasing faster at the outer surface than inside the specimens (Figs. 5.29 and 5.30). Stage II starts when the surface layer is totally dry and, hence, the drying front begins receding into the specimen (Figs. 5.29 and 5.30).

The main difference between the two specimens is that, as expected, the unpainted specimen dries faster. The difference is particularly significant during drying stage I, when a clearly lower drying rate occurs for the painted than for the unpainted specimen (Figs. 5.30 and 5.31). As a consequence, the drying front recedes earlier into the unpainted specimen (Fig. 5.29) and, hence: (i) the surface of the painted specimen stays wet longer; (ii) a higher amount of moisture is eliminated by surface evaporation on the painted specimen.

Reduction of the evaporation rate is likely to be due to the paint reducing the ESE (effective surface of evaporation) on the specimen. This arises probably because the paint blocks part of the surface pores of the stone. However, it is also possible that the water/paint contact angle is lower than the water/mortar contact angle. Therefore, a lower curvature of the menisci at the surface could have also contributed to reducing the ESE, as discussed in section 4.5.4. This kind of effect was suggested by Rousset-Tournier (2001) to explain why Stage I drying rate was higher for saturated stones than for a free water surface.

Another interesting observation is that, also here, the drying front is not a sharp front. Indeed, Fig. 5.30 shows that the surface layer starts very early on loosing moisture faster than the rest of the specimen, despite the drying front only receding completely into the material some hours later. As seen in Figs. 5.29 and 5.30, the surface begins drying faster in less than 1h (unpainted specimen) or after around 2h (painted specimen) but the front only recedes completely into the material after 2h up to around 4h or after 8h to 10h, respectively.

This raises the question of whether, in the case of a diffuse drying front, the total retreat of the front into the material corresponds exactly, as currently admitted (Mertz 1991, for instance), to the critical point where the overall drying curve changes from linear into an asymptotic curve (Fig. 4.6). Indeed, Fig. 5.31 suggests that this transition point may occur, particularly in the case of the unpainted specimen which gave rise to a more diffuse front, some time before the total retreat of the drying front into the material.

In both specimens, as soon as the moisture content begins decreasing at the surface, an approximately linear variation of the moisture content arises between the surface and the middle of the specimens, which corresponds to a markedly diffuse drying front. In the latter moments of the experiment, a drying front width of around 13 mm or 9 mm (see green lines in Fig. 5.30) is observed for the unpainted and painted specimens, respectively. This width difference is coherent because the slower the drying, the sharper the drying front. Indeed, a lower drying offset is expected to exist among pores of different sizes when evaporation is slower, as it happens with the painted specimen, because the liquid has more time to flow to the surface.

5.8 Drying of hydrophobic agent containing plaster (set 4)

5.8.1 - Introduction

Hydrophobic agents are often added to plastering/rendering mortars aiming at preventing dampness or salt damage. Typically, small amounts of such additives are used to reduce the capillarity coefficient of these mortars without rendering them into fully hydrophobic mortars. The current conviction is that a lower capillarity coefficient will make it more difficult for liquid solutions to reach the outer surface of walls and, thereby, surface damage will be reduced.

However, there is field evidence that surface damage, dampness included, is in certain cases enhanced rather than reduced when hydrophobic agent containing plasters or renders are used (Fig. 5.2).

Drying experiments monitored by a 2D MRI technique were therefore performed on a traditional lime plaster that contained a hydrophobic agent. The objective of these experiments was to evaluate the influence of the additive on drying and salt deposition processes.

5.8.2 - Materials and methods

The drying experiments were similar to those presented in Chapter 4 where the 2D MRI technique described in Annex I was used on specimens composed of a lime plaster on a composite substrate of lime mortar and stone. But here, lime plaster H was used instead of plaster L.

The composition and execution technique of these two plasters is similar in all respects except for the hydrophobic agent in plaster H, which was added during batching in the proportion recommended by the producer (Table 5.1). Therefore, the influence of the additive can be evaluated by comparing the behaviour of the two types of specimens. As seen in Fig. 5.5, the additive induced a certain lowering of the capillarity coefficient of mortar H by comparison to mortar L.

The experiments were performed on specimens filled with either pure water (specimens DH3 and MH3) or 3 molal NaCl solution (specimen DH1). Drying was monitored for 20 hours for pure water and 50 hours for the NaCl solution.

Also gravimetric drying experiments, similar to those described in section 5.4, were performed on several DL and DH specimens, as well as on similar specimens where stone M was used instead of stone D (ML and MH specimens). A first series of gravimetric experiments was carried out at around the two months of age of the specimens, that is, approximately three months before the MRI-monitored experiments were performed. Three speciments of each kind were tested, including those later subjected to the MRI-monitored experiments. Approximately three months after these MRI-monitored drying experiments were performed, a new series of gravimetric drying experiments was carried out on four specimens, three of which (DL5, DH10 and MH10) had never been wetted.

5.8.3 - Results

Selected MRI snapshots of specimen DH3 during drying with pure water are presented in Fig. 5.32. Snapshots of specimen DL1 (Chapter 4) under similar drying conditions are also included to make it easier a comparison between plasters with (DH3) and without (DL1) hydrophobic additive. Snapshots that correspond to approximately similar moisture distributions in the two specimens, though to different moments of drying, were chosen. In Fig. 5.33 the 1D distribution profiles at cross-sections CS1 and CS2 (Fig. 4.10) of the two specimens are presented.







Fig. 5.33 – Drying with pure water: 1D moisture profiles at cross-sections CS1 and CS2 of specimens DH3 and DL1. MRI signal is given in arbitrary units (a.u.). The grey shadow areas indicate the possible location of the physical limits of the test specimens (the drying surface is always on the left).

The overall drying curves are shown in Fig. 5.34. This figure includes not only the drying curve of specimen DH3 that concerns the drying experiment presented in Figs. 5.32 and 5.33, but also the drying curves of two other specimens: MH3 (pure water) and DH1 (NaCl solution). Drying curves of specimens with similar composition but where plaster L is used instead of plaster H are also included.

Each point on a drying curve corresponds to the sum of all the values obtained across the two corresponding 1D profiles (similar to those in Fig. 5.33). Normalized drying curves are presented in this case to make it easier to compare the different specimens.



Fig. 5.34 – Normalized drying curves

The results of the gravimetric drying experiments performed on various specimens at the two or eight months of age are presented in Fig. 5.35.



Fig. 5.35 – Gravimetric drying experiments on water-filled specimens composed of non-additivated plaster L or additivated plaster H. The experiments were performed at the two (graph on the left) or eight (graph on the right) months of age.

5.8.4 - Discussion on set 4

With pure water, Fig. 5.32 shows that the overall drying behaviour of specimens DH3 and DL1 leads to equivalent situations which, nonetheless, correspond to different moments. In fact, drying is slower for the hydrophobic plaster. Fig. 5.33 shows that the evaporation rate during drying Stage I is indeed lower for specimen DH3 than for specimen DL1. As a result of this slower drying process, the drying front receded at a later moment in specimen DH3. This behaviour may account for the fact that in real walls dampness and surface decay are sometimes worsened when hydrophobic additives are used in plastering or rendering mortars (Fig. 5.2).

A higher offset seems to exist in specimen DH, particularly in the early moments, between the moisture content in the plaster and in the substrate. This suggests that the hydrophobic agent reduces the moisture content in the plaster. Indeed, Fig. 5.5 shows that not only is the capillarity coefficient reduced by the use of this additive but also capillary saturation seems to be attained at lower moisture content.

Hydrophobic products used on porous building materials are substances that possess a polar and a non-polar end. The polar end is adsorbed into the polar surfaces of the pores thereby giving non-polar characteristic to these surfaces. Thus, the contact angle between water and such surfaces is higher than 90°. In pores, a convex meniscus forms and there is no capillary suction. Mass hydrophobic additives used in mortars are liquid or powder products. It is known that the higher the quantity of product added, the lower the capillarity coefficient of the material. It is reasonable to assume that this arises from the hydrophobation of an increasing part of the porosity.

Therefore, a part of the porosity in plaster H was probably unavailable for capillary transport. This is consistent with the behaviour differences between specimens DH3 and DL1:

- Mortar L and mortar H have similar pore size distributions (Fig. 5.7). Yet, in contrast to specimen DL1, the moisture content in the plaster in specimen DH3 is always somewhat lower than that in the underlying mortar L (Figs. 5.32 and 5.33).
- The slower drying of specimen DH3 is likely to arise mostly from that lower effective volume of capillary active pores through which a practically similar amount of water has to be eliminated. A lower liquid flux is accompanied by a lower evaporation rate because less liquid is available for evaporation at the wet front.
- It is not known if there is also a reduction of the vapour flux in hydrophobic pores. Indeed, in non-hydrophobic pores, the presence of liquid water can enhance vapour transport (Rose 1963) as the so-called "liquid assisted vapour transfer" (Fig. 4.5c).

The longer drying stage I of specimen DH3 in comparison to specimen DL1 is not very clear in the overall drying curves in Fig. 5.34. This may occur because, since the specimens are not homogenous, the drying front recedes unevenly throughout the surface, as roughly suggested by Fig. 5.32. This is consistent with the fact that in the 1D profiles (Fig. 5.33) no effective receding of the drying front into the material is observed. Indeed, the drying front seems to recede somewhat later at the central parts to which these profiles correspond (Fig. 1.10) than at the edges of the specimen (Fig. 5.32). A second cause can arise from the drying front being diffuse rather than sharp. Hence, the effective surface of evaporation and, thus, the drying rate of the specimen, can start decreasing before the drying front recedes completely into the material, as discussed in the previous section.
Only one of the three MRI-monitored drying experiments performed on plaster H specimens has been discussed up to now. Indeed, the two other drying experiments on specimens MH3 (pure water) or DH1 (NaCl solution) were not conclusive. In these experiments, the additivated and non-additivated plaster showed similar drying behaviour, as seen in Fig. 5.34. This occurred despite lower evaporation rates were measured for all the specimens with plaster H in the first gravimetric experiments (Fig. 5.35, on the left). To clarify these unexpected results, new gravimetric drying experiments were performed some months later. As seen in Fig. 5.35 (on the right), the hydrophobic additive had completely lost its effect, despite most of the specimens never having been wetted. Therefore, the similar drying behaviour of specimens MH3 and ML1 with pure water or DH1 and DL1 with NaCl solution was most likely due to an absence of hydrophobic effect. Further research on this loss of hydrophobic effect could not be carried out within this thesis due to time limitations and also because it was out of the scope of the original working plan.

5.9 Crystallization tests on painted stone (set 5)

5.9.1 - Introduction

Crystallization tests are typically performed on one-material specimens. Yet, in reality, walls frequently include a final paint layer. Furthermore, salt decay problems in these walls often consist essentially of surface damage due to paint peeling or blistering.

In this section, NaCl and Na_2SO_4 crystallization tests are presented, which were carried out on painted and unpainted stone specimens with the main objectives of:

- comparing the damage caused by each type of salt;
- evaluating the performance of the painted and unpainted material.

5.9.2 - Materials and methods

Two distinct crystallization tests were performed on stone D specimens either unpainted or painted with common acrylic emulsion paint.

Experiment 1

The specimens were similar to those in set 3, although of larger flat dimensions (160 mm x 160 mm). The lateral faces of the specimens were sealed with silicon mastic. Later, it was found that this type of silicon product can migrate into the pore network and cause hydrophobization of stone, mortar or brick materials. Here, this type of effect indeed arose in a peripheral band which attained a maximum width of 5 mm in some specimens. Yet, due to the large size of these specimens, there were no major consequences.

Crystallization tests by wet/dry cycling were performed, using pure water, 0.4 molal NaCl solution or 0.4 molal Na_2SO_4 solution. Two specimens of each kind were used with either salt solution. For pure water, only one specimen of each kind was used. After oven drying the specimens at 60°C for 24 hours, four cycles were carried out as follows:

- wetting the specimens were immersed in the solution to a depth of around 5mm from their bottom faces for 2 h;
- the bottom faces were sealed with polyethylene film;
- drying the specimens were dried in a ventilated oven at 60°C for 24 h;
- cooling the specimens cooled for 22h in a conditioned room with a low air velocity at 20°C and 65% RH;
- the specimens were observed and any damage registered;
- the wet-dry cycle ended.

Experiment 2

A second test was performed only on painted stone. Smaller 100 mm x 100 mm specimens laterally sealed with epoxy resin were used in this case.

Wet/dry cycles were performed, using 0.2 molal NaCl solution or 0.1 molal Na_2SO_4 solution. Two specimens were used with each salt solution. Three cycles were carried out after oven drying the specimens at 60°C for 24 hours. The cycles were similar to those in the previous experiment with the difference that the cooling and drying periods were reduced from 22 to 2 hours and from 24 to 22 hours, respectively. These changes allowed total cycle duration of 24h. Also, observation and registration of damage was now carried out at the end of each absorption, drying or cooling period.

5.9.3 - Results

Experiment 1

In the first experiment, extensive damage was observed on all painted specimens already at the end of the second cycle. The paint blistered with both salt solutions, although the blisters were slightly larger for Na_2SO_4 than for NaCl. At the end of the fourth cycle, large areas of the paint had become completely detached from the substrate (Fig. 5.36), especially in the case of Na_2SO_4 .



Fig. 5.36 - Damage on the acrylic paint at the end of the first crystallization experiment

The unpainted specimens developed efflorescence both for sodium sulfate and for sodium chloride. As seen (Fig. 5.37, left), sodium chloride efflorescence was much more abundant, forming a thick crust of salt that eventually blistered. Yet, after efflorescence removal, the surface of the stone was eroded in the case of sodium sulfate but practically undamaged in the case of sodium chloride (Fig. 5.37, right).



Fig. 5.37 - Damage on the unpainted stone specimens at the end of the first crystallization experiment: efflorescence (left), cleaned stone surface (right)

With pure water no damage arose neither for the painted nor for the unpainted specimens.

Experiment 2

Extensive damage was also achieved in the second crystallization experiment with both salts. Damage arose during the second or third cycle for sodium sulfate or sodium chloride, respectively (Fig. 5.38). It is interesting to note that sodium sulfate damage occurred during the wetting period, while sodium chloride damage arose during the drying period. By perforating the blistered paint layer it could be seen that there was no salient efflorescence that could have pushed the paint. The surface of the stone was almost clean in the case of sodium sulfate and for sodium chloride only a thin layer of salt was visible.

Liquid water was systematically observed on the painted surfaces during wetting of the specimens. This indicates that the specimens were able to saturate during the two-hour immersion. That is consistent with the results of the capillary absorption test (Fig. 5.5) were capillary saturation of stone D was attained after one hour.



Fig. 5.38 - Damage to the acrylic paint during the second crystallization experiment: for sodium sulfate, after one hour soaking in the second cycle (left); for sodium chloride, at the end of the third cycle drying (right). The paint layers were intentionally perforated to allow observing the stone surface.

5.9.4 - Discussion on set 5

To the unpainted stone, sodium chloride caused much less material damage than sodium sulfate, as seen by the results of the first experiment (Fig. 5.37). As in many other crystallization tests (Goudie and Viles 1997, for instance), sodium chloride formed mostly harmless efflorescence while sodium sulfate crystallized mainly within the pores thereby eroding the stone.

Yet, both salts caused significant damage to the painted specimens, though the damage mechanism is not completely clear. Indeed, there was no efflorescence beneath the paint layer that could have pushed it, which contradicts the hypothesis put forward in a previous article (Gonçalves 2003a). But, regardless of the actual damage mechanism, it is clear that both sodium sulfate and sodium chloride can cause severe damage to painted specimens even in current crystallization tests where only sodium sulfate is able to deteriorate the unpainted material.

Because paint damage is one of the most frequent salt decay patterns in buildings, it will be important to clarify the actual damage mechanisms in future work. The main hypotheses are the following:

- (i) Vapour pressure Salt deposition at the paint/plaster interface can weaken the adherence of the paint to the substrate or even create a thin air layer at the interface. The paint hinders vapour transport, as seen in section 5.8. Hence, it is possible that, in subsequent cycles, vapour pressure is able to push the poorly adherent paint layer which then blisters and eventually detaches completely. Indeed, no damage arose with pure water (experiment 1) and salt deposition was observed at the paint/plaster interface of the damaged specimens (experiment 2).
- (ii) Crystallization pressure In the second experiment, the painted surfaces become wet during the wetting periods. Hence, some salt probably crystallized in the paint layer itself. Therefore, it is possible that crystallization pressure causing dilation of the thin, poorly adherent and elastic coating. Indeed, in this experiment, sodium chloride damage arises during the drying period, when halite is expected to crystallize mostly. In contrast, damage by sodium sulfate arises during the wetting period, hence, it may be due to mirabilite crystallization at the contact of the fresh salt solution with thenardite crystals deposited in the previous drying phase.

5.10 Discussion

5.10.1 - Salt distribution

In the case of evaporative crystallization, soluble salts crystallize at the drying front. The depth and width of the layer at which they deposit in porous materials depends on the position and width of the drying front. The distribution of salt is, therefore, controlled by all factors that influence evaporation or liquid movement to the drying front, as early postulated by Schaffer (1932).

The overall position of the drying front results, at each moment, from a global balance between the incoming liquid flux and the outgoing evaporation flux. The lower the liquid/evaporation flux ratio, the deeper in the wall is the drying front located and, thus, soluble salts deposited.

The width of that front also conditions the distribution of salt, as argued in section 5.6.4.2. When a diffuse front occurs, the solubility of the salts is relevant: (i) the lower this solubility, the earlier the salts will crystallize as they travel across the front; (ii) for salts that tend to supersaturate or possess several crystal phases with distinct solubility, such as Na_2SO_4 , a more even distribution of salt can arise across the drying front. Indeed, a very diffuse drying front was observed in MEP plaster in set 2. And both in set 1 and 2 a deeper and more even deposition of Na_2SO_4 than NaCl occurred, namely, in MEP plaster.

There are some differences between the distributions of Na_2SO_4 across MEP plaster in the two sets: in set 1 more salt crystallized at the plaster middle-thickness (Table 5.4) while in set 2 the salt accumulated preferentially next to the plaster/substrate interface (Fig. 5.26). Further, drying was globally slower in set 1 than in set 2 (Fig. 5.12 and 5.21). Yet, a more advanced position of the drying front during set 1 is not probably the reason, because the two NaCl distributions are reasonably similar (Table 5.4 and Fig. 5.25). These small differences between the distributions of Na_2SO_4 in the two sets are not easy to explain. It might be that the repeated contaminations or temperature cycles in set 1 had some influence. Indeed, temperature-induced crystallization of mirabilite or crystallization of mirabilite at the contact of the salt solution with pre-existing thenardite crystals may have occurred in set 1. But a distinct metastable behaviour of sodium sulfate, as regards supersaturation or heptahydrate precipitation, is equally possible.

The width of a drying front depends on the homogeneity of the pore network. In heterogeneous pore networks (as regards both pore suction and size) the offset between the positions of the local drying fronts in the different pore families can be significant. Hence, a diffuse instead of a sharp front might arise.

Furthermore, all factors that influence the liquid flux or the evaporation rate can, in principle, alter the width of the drying front. For instance, the slower drying of the painted specimen in set 3 gave rise to a sharper drying front (Fig. 5.30). Indeed, a smaller drying offset exists among pores of different sizes when evaporation is slower because the liquid has more time to flow to the evaporation front.

Therefore, salt distribution in plasters and renders depends on the (vapour and liquid) moisture transport properties of the material but may also be influenced by external factors such as:

- i) Those that influence the evaporation rate for example: nature (and concentration) of the contaminant solution, environmental conditions and presence and type of paint layer.
- ii) Those that condition liquid transport, for instance: moisture content in the materials and suction properties of the substrate.

5.10.2 - Plasters and renders

In this chapter, the behaviour of plasters and renders in relation to soluble salt crystallization was described in terms of their salt-accumulation behaviour, damage patterns and durability. Efflorescence was considered as damage for the salt-accumulating but not for the salt-transporting plasters.

As seen in set 1, a general comparison of the behaviour of different plasters in relation to salt crystallization is not possible because it may change with distinct boundary conditions. Indeed, only Parlumière presented reasonably constant (salt-accumulating) behaviour under different testing conditions. Further, the durability of all plasters varied significantly with the type of contaminant solution, presence of paint or type of substrate material.

Soluble salts slow down evaporation. This occurs because non-volatile solutes lower the vapour pressure of liquids and also probably because of the hindering effect of salt crystals (as seen in set 1, the drying rate of the salt-loaded specimens decreased from cycle to cycle). Hence, the outer surface stays wet longer when salts are present. The lower the vapour pressure of the salt solution, which depends on the type of salt and concentration of the solution, the closer to this surface salts deposit during evaporative processes.

Yet, different plasters are affected differently by the presence of salts. In set 1, the decrease in the drying rate from cycle to cycle was more severe in the case of salt-transporting (LNEC on brick) than of salt-accumulating behaviour (LNEC on lime mortar, MEP and Parlumière on brick). Further, some systems showed similar salt-accumulating behaviour with NaCl and Na₂SO₄ while others did not. The most extreme change occurred with painted MEP which worked as an accumulating or as a transporting plaster with Na₂SO₄ and NaCl, respectively. Also, although all systems dried more slowly when salts were present, different drying speed rankings arose for pure water and the salt solutions. Since pure water corresponds to an infinitely dilute solution, this suggests that solution concentration differences may affect the behaviour of distinct plasters differently.

Paints can also hinder evaporation, hence reducing the ratio evaporation flux / liquid flux and causing a more superficial deposition of salt and a longer presence of moisture at the surface. This was clearly shown by the MRI images in set 3 drying experiments. Also in set 1 drying tended to be slowed down by the presence of the paint layer both for pure water and for the salt solutions. However, the influence of the paint varied for different plasters and salts. For instance, with NaCl, the paint transformed MEP from an accumulating to a transporting plaster. But this did not occur for MEP with Na_2SO_4 nor, with any of the salts, for Parlumière or LNEC on lime mortar.

The influence of paints is expected to be different in the case of salt-accumulating or salttransporting plasters. For salt-accumulating plasters, serious damage may arise from the use of paints because, when drying is slowed down, salt solutions can reach the surface and, hence, completely corrupt the fundamental working principle of the plaster. In contrast, for salt-transporting plasters, paints that hinder drying may tend to increase salt-induced dampness and efflorescence but they can, at the same time, reduce material damage by salt crystallization. This was clearly observed in set 1 tests on LNEC/brick with NaCI: surface damage arose earlier on the painted specimens but it was less severe (only efflorescence and further rupture of the paint layer) than on the unpainted specimens (where longitudinal cracking occurred).

Mass hydrophobic additives, when used in reasonably low amounts that do not render a traditional plaster into a fully hydrophobic plaster, cause reduction of the liquid flux because a part of the capillaries becomes unavailable for liquid transport. Hence, the evaporation rate is also reduced because less liquid is available at the wet front per unit time. Therefore, drying is prolonged. In the case of specimen DH in set 4, which was initially saturated, this led to a longer drying Stage I.

The substrate material can strongly condition the liquid flux into the plaster, hence, the evaporation/liquid flux ratio, thus, the behaviour of plasters and renders in relation to salt crystallization. As seen in set 1, LNEC behaved as a transporting plaster on brick substrate and as an accumulating plaster on lime mortar substrate. This indicates that LNEC is able to draw a higher amount of moisture per unit time from the brick than from the lime mortar. This probably occurs because the lime mortar possesses a higher amount of small pores than the brick (Fig. 5.7). Hence, a higher percentage of the pores in LNEC plaster is able to draw water from the brick than from the lime mortar substrate. Indeed, the liquid flux depends on the relation between the capillary suction of the two materials at different moisture contents. There is also a possibility that a poorer adherence existed to the lime mortar substrate thereby inducing a higher resistance to liquid transport. Indeed, resistance to liquid transport may vary at the interface between different materials.

The saturation condition is also relevant as regards the behaviour of porous materials to salt crystallization. As seen in set 3, when MEP specimens were fully saturated, most of the moisture was eliminated by surface evaporation and, hence, a markedly transporting behaviour arose. In contrast, salt-accumulating behaviour occurred when only the substrate was saturated. In any case, absorption/drying hysteresis prevents an immediate retreat of the wet front at decrease of the moisture content in the material. This suggests that strong hysteresis can be positive for plasters intended to be salt-transporting. For salt-accumulating plasters it may also contribute to a longer stability of the drying front, thus, to a lower risk of salt being deposited on or in the substrate as a result of diminishment of the liquid flux. However, if the boundary conditions are such that the wet front is able to reach the surface, a strong hysteresis will worsen surface damage in salt-accumulating plasters.

In real walls rising damp represents the general case and, unlike in laboratory unidirectional drying tests on small specimens: (i) moisture transport is three-dimensional or can, at the best, be considered bidirectional when sufficiently far from the lateral limits of the wall; (ii) permanent liquid supply fluxes tipically exist during more or less prolonged periods; (iii) the moisture content in the wall varies (decreases) with height. It is common knowledge that lowering of the evaporation / liquid flux ratio, for instance due to temperature drop or RH rise, increases the height of capillary rise in walls. Hence, all the above mentioned factors that lower the ratio evaporation flux / liquid flux not only induce, as argued, a forward displacement of the drying front at each height, but also an upward increase in the height of capillary rise. Indeed, they tend to cause a general enlargement in the moist or salt decayed area.

In the MRI-monitored experiments noticeably diffuse drying fronts were often observed. Indeed, a sharp front arose only, in set 2 and in Chapter 4 experiments, at the surface of the specimens totally saturated with salt solution when the duration of the experiment was perhaps too short to allow receding of the drying front. In the other cases, markedly diffuse drying fronts arose. Although drying in these experiments was probably much faster than in most real situations, it is probable that for low liquid supply fluxes or high evaporation rates (for instance, due to wind) diffuse wet fronts will also arise in building masonry.

Drying curves, normally obtained by means of gravimetric drying experiments, are currently used to characterize building materials. The critical moisture content in those curves (Fig. 4.6) corresponds to the moment when the drying rate starts decreasing over time. This point is expected to represent the transition from Stage I (when the drying front is located at the surface) to Stage II (when the drying front is located in the material). However, if the drying front is not sharp, transition from Stage I to Stage II is not immediate. The drying front only recedes completely into the material after a transitional period where the porosity involved in transporting the liquid to the surface decreases progressively. The more diffuse a drying front, the longer that transitional period, as seen in set 3 (Fig. 5.31). This suggests that drying curves, either gravimetric or obtained by other methods such as MRI, may not be totally reliable for evaluating drying of porous building materials. Indeed, for a sharper front, the critical moisture content may indeed correspond to the moment when the surface layer becomes dry. But there is no guaranty that the same will happen for a more diffuse front, where that point may simply correspond to a decrease in the water-filled surface porosity.

Moreover, the sharpness of the drying front does not depend solely on the material properties. A slower drying rate, for instance due to differences in the environmental conditions or presence of a paint layer, can give rise to a sharper front in the same material and for the same liquid. This misleading effect is suggested, for instance, when comparing the behaviour of the painted and unpainted specimens in set 3. While the critical point in the overall drying curve of the painted specimen is close to that when the drying front totally recedes into the material, it is not clear that the same happens for the unpainted specimen (Fig. 5.31).

EN standard 998-1 (CEN, 2003) defines requirements for the so-called renovation mortars that concern capillary absorption and vapour permeability (Table 2.1). Renovation mortars are here defined as "plastering/rendering mortars to be used on moist salt-loaded walls", which is a rather insufficient specification. Indeed, nothing is said about the working principle which, from the end-users' point of view, is perhaps the most fundamental feature. Although it was not an objective of this thesis to verify the fulfilment of standard requirements such as these, the results obtained in the several sets of tests indicate that material properties are insufficient for predicting the behaviour of plasters and renders in relation to salt crystallization. Indeed, only in the case of salt-blocking (fully hydrophobic) or moisture-sealing plasters can constant behaviour be expected for different boundary conditions. Hence, for transporting or accumulating plasters a probabilistic approach based on field data would perhaps be more adequate, even if not easy to achieve. Another possible way is the use of numerical computational models that allow evaluating the performance achieved under varied conditions. However, further knowledge of salt decay mechanisms is still necessary, namely of the processes by which salt crystals hinder drying, in order to develop adequate models, as also concluded by Huinink and Pel (2003).

5.10.3 - Sodium chloride and sodium sulfate

Although there is field evidence of its frequent involvement in salt decay processes, sodium chloride is often, in contrast to sodium sulfate, almost totally or totally harmless in laboratory tests.

The different behaviour of sodium chloride and sodium sulfate probably lies, to a significant extent, in differences between the crystallization processes of the two salts, namely, in the capability of sodium sulfate for generating highly supersaturated solutions, hence, strong crystallization pressures (Tsui *et al.* 2003) or in the faster crystallization kinetics of mirabilite in comparison to halite (Rodriguez-Navarro and Doehne 1999). Further, in contrast to sodium chloride, the solubility of sodium sulfate varies with temperature (Figs. 2.6 and 2.8), hence, temperature-induced crystallization may take place. Indeed, laboratory crystallization tests often include temperature cycling, frequently at very high temperatures. It is also possible that the enthalpy change associated to the crystallization process itself (which is endothermic in both cases) is, due to the sensitivity of this salt to temperature changes, relevant for sodium sulfate. Mirabilite crystallization may also occur readily at the contact of a sodium sulfate solution with pre-existing thenardite crystals (Chatterji and Jensen 1989).

Another point may be that mixtures of salts are usually present in buildings while in laboratory tests single salts are normally used. And the properties of soluble salts may change in the presence of other salts (Price and Brimblecombe 1994). In fact, a combination of different salts can be more harmful than each of them in isolation (Charola 2000).

Further, specific damage patterns such as alveolization (Fig. 2.3) are not reproduced in current crystallization tests.

The experimental work performed in this thesis indicates that other differences between the properties of sodium chloride and sodium sulfate, as well as differences between ordinary laboratory and field conditions may also contribute to these behaviour discrepancies.

Laboratory tests are normally carried out: (i) on specimens made of a single material; (ii) on fully-saturated specimens; (iii) using very concentrated solutions. These features can account for lower crystallization damage by sodium chloride than by sodium sulfate because: (i) the vapour pressure of similarly concentrated salt solutions and, particularly, of saturated solutions is lower for sodium chloride; (ii) at ordinary temperatures, the solubility of the possible crystal phases of sodium sulfate (thenardite, heptahydrate and mirabilite) is lower than the solubility of halite, the crystal phase of sodium chloride.

Faster drying typically arises for sodium sulfate than for sodium chloride, namely, because of the higher vapour pressure of similarly concentrated and, particularly, of saturated solutions. Accordingly, sodium sulfate tends to crystallize deeper in the material because: (i) the drying front is more deeply located; (ii) the drying front is more diffuse; (iii) due to its lower solubility, sodium sulfate tends to crystallize earlier in the drying front. In contrast, for sodium chloride, a more superficial and sharper drying front will arise thereby inducing a more superficial deposition of salt. This feature is stressed when very concentrated salt solutions are used, as often happens in laboratory tests. For saturated solutions, for instance, the vapour pressure difference is huge: at 20°C, the RH_{eq}^{sat} is of 75% for sodium chloride and 95.6% for sodium sulfate. Since fully saturated specimens are the rule in laboratory tests, drying stage I will be

much longer for sodium chloride, as seen in Chapter 4. Hence, a much higher percentage of sodium chloride than of sodium sulfate will crystallize as harmless efflorescence. That was clearly observed in the crystallization experiments of set 5 on unpainted stone.

In contrast, when the testing conditions are such that the drying front is inside the material during the entire drying process, sodium chloride can cause even more severe damage than sodium sulfate, as seen in set 1. This is likely to result, to a significant extent, from a more localized deposition of sodium chloride. As observed in sets 1 and 2, sodium sulfate tends to be spread in a larger volume probably because of its ability for supersaturating, for precipitating at different concentrations and, in some cases, for crystallizing due to temperature changes or at the contact between the salt solution and pre-existing crystals.

Unpainted specimens are normally used in laboratory tests. In contrast, painted surfaces are common in buildings. Further, salt decay patterns in those buildings often consist simply in peeling of the paint layer and efflorescence. This difference can probably account for part of the discrepancy between the in-lab ineffectiveness and on-field effectiveness of sodium chloride. As seen in set 5, only sodium sulfate was able to deteriorate the unpainted stone. Yet, both sodium sulfate and sodium chloride were able to cause severe damage to the painted specimens.

In set 5 crystallization tests the paints blistered and peeled without having been pushed by efflorescence. This sort of damage is often seen in buildings (Fig. 5.39). But paint damage can also arise due to subflorescence pressure (Fig. 5.40). This second kind of damage is normally associated to high liquid fluxes (for instance, at the base of walls) and low evaporation rates (for instance, in poorly ventilated rooms). It is not known whether the two damage mechanisms are predominantly related to particular types of salts. Nevertheless, it is sodium chloride able reasonable to suppose that is to cause significant subflorescence-induced damage, perhaps even more than sodium sulfate. Because of the high tendency of sodium chloride to crystallize on the surface rather than on the pores of building materials, a large amount of salt can probably crystallize beneath, for instance, vapour permeable hydrophobic paint layers. Indeed, as seen in Fig. 5.1, halite crystals are even able to pull off a 1 cm thick hydrophobic brick layer.



Fig. 5.39 – Paint damage arising without the presence of efflorescence: interior of Ajuda Palace, Lisbon (left) and exterior of building in Almada, Portugal (right).



Fig. 5.40 – Paint damage by subflorescence pressure: Alhos-Vedros tide-mill (left) and interior of Despacho House, Pereira, Portugal (right).

5.10.4 - Development of an accelerated-ageing test

It is not always easy to understand the criteria underlying salt crystallization protocols. This was well expressed by Goudie and Viles (1997) when commenting on a certain crystallization test:

We might ask questions about the basis for the concentration of sodium sulfate, why sodium sulfate was used, why the humidity is not carefully controlled, why such an extreme drying temperature is used, why the soaking phase has the length it has. Similar questions might be asked of [other methods].

Ultimately, it is not simple to design crystallization tests. Test variables should represent, as closely as possible, a complex, dynamic, changeable and multiple reality from which there is often not enough field information available. And it is not straightforward to define these variables to produce short-duration (stronger) action on small specimens than reasonably simulates long-duration (weaker) action on buildings. As stated by Knöfel *et al.* (1987) about weathering tests in general, intensified experimental conditions lead to more destructive processes, which take place only under more severe conditions, rather than just to a time lapse of weathering processes.

Indeed, there is a certain contradiction between two of the main aims of salt crystallization tests: (i) simulating reality; (ii) speeding up damage by comparison to reality. The key points are that:

- i) Salt decay dynamics influences the type of decay. As seen in this thesis, all factors that influence evaporation or liquid transport condition the position and width of the drying front, hence, the type of decay. Therefore, when damage is accelerated, by harshening the environmental conditions or by using very concentrated solutions or high moisture contents, distinct decay process and features will most likely arise.
- ii) This influence varies for different materials and boundary conditions. For example, in set 1, different drying speed rankings arose for pure water and the salt solutions, which suggest that the concentration of salt solutions has a different influence on the behaviour of different materials.

Salt crystallization tests can be used to compare the influence of different factors on salt decay processes, as seen in this thesis, or to investigate salt damage mechanisms in different materials. Yet, for the above mentioned reasons, they can hardly be used as general performance tests for building materials. This conclusion contradicts Schaffer's (1932) statement that "crystallization tests (...) tend to reproduce that type of weathering which the material suffers under actual exposure to the weather". Indeed, the influence of the experimental conditions on the type of decay is such that simulating and, at the same time, accelerating the process is probably impossible.

5.10.5 - Restoration interventions

End-users complain that salt decay in buildings, particularly efflorescence and material damage such as peeling of paints or surface sanding of plasters and renders, is often worsened after restorations.

Reduction of the ratio evaporation/liquid flux by the use of new materials is one possible cause. Indeed, when that ratio is lowered, a forward displacement of the wet front arises for a given liquid flux. This occurs when materials of lower vapour permeability than the original ones are used, for instance, emulsion paints instead of pure lime washes or cement-based instead of lime-based plasters and renders.

In walls with rising damp, the moisture content decreases with height. The water rises up to the height where the liquid supply flux from the ground is balanced by the evaporation flux through the surface of the wall. Therefore, hindering of evaporation, for instance by the use of paints, may have two consequences: (i) at each height, the drying front moves closer to the surface; (ii) there is an increase in the overall height of capillary rise (this reasoning is applicable also to the lateral enlargement of the area affected by salt solutions). Therefore, enlargement of the moist or salt-damaged area or damage to adjacent elements can arise from the use of materials with low vapour permeability. One typical situation is that when moist salt-damaged walls in the interior of rooms are repeatedly painted. Each new paint layer will further reduce the evaporation rate. Hence, the damaged surface area is progressively extended.

Reduction of the evaporation flux and further worsening of damage might also arise from a reduction in the liquid flux. This occurs when the lowering of liquid transport is not enough to, under certain environmental conditions, prevent the liquid from reaching the surface of the wall. In that case, (surface) evaporation occurs at a slower rate, hence, the damaged area increases. This is typical of the so-called "slow-transporting" plasters and renders, for example, traditional plasters with small amounts of hydrophobic additives. But a similar effect may arise also with plasters and renders that generally work as fast-transporting if there is:

- i) a deficient bond to the substrate: lower liquid flux into the plaster may occur as a result of poor connectivity of the pores at the interface with the substrate;
- ii) poor compatibility between the pore networks of the plaster and substrate: lower liquid flux to the surface arises because only a reduced percentage of the (small) pores in the plaster is able to draw moisture from the substrate.

Surface damage on salt-accumulating plasters may occur, as explained, from the use of paints of low vapour permeability or from repaintings. Also, as seen in set 2, if solutions are allowed to reach the surface, by whichever conditions that lower the evaporation rate, surface evaporation can persist long after that event has ceased due to absorption/drying hysteresis of the plaster. This can happen also when the plaster or render is externally wetted, for instance, by dew point condensation or rain and, hence, liquid continuity up to the surface is enabled. For this reason, the use of a hydrophobic surface layer on salt-accumulating systems seems recommendable for accumulating plasters and unavoidable for accumulating renders.

Liquid continuity up to the surface can, theoretically, be established also by the water in fresh plastering/rendering mortars. It is reasonable to suppose that surface evaporation may afterwards persist, during or even after hardening, in plasters/renders with strong hysteretic behaviour or under low evaporation conditions (for example, during cold or wet seasons, in poorly ventilated rooms or when paints are applied before the plaster is completely dry).

Conversely, problems may arise in fast-transporting plasters and renders from an acceleration of the evaporation rate. It is known that when, for instance, dehumidifiers are used aiming at avoiding efflorescence and dampness on these materials, a somewhat deeper or more diffuse drying front may occur. Hence, a higher amount of salt may crystallize in the pores close to the surface and induce material damage instead of harmless efflorescence.

5.10.6 - Adequacy of plasters and renders

In monuments conservation, avoiding surface damage is normally not the only (and often not the main) objective. Preservation of the underlying masonry and adjacent elements is also frequently necessary. Indeed, different requirements arise in distinct situations:

- Health requirements are very relevant for living spaces, museums or rooms where perishable goods are stored, such as paper, food, etc. In this case, surface damage (dampness, efflorescence or material damage) is hardly acceptable.
- Protection of the substrate is important for buildings of high historic value and for weak masonry.
- Sometimes there are artistic elements in continuity to the plastered/rendered area. In this case, it is necessary to avoid that salt solutions are diverted into them.

Therefore, different plasters and renders may be adequate for distinct situations. Furthermore, a wide range of situations can account for an increase in damage after restoration interventions, as seen in the previous section for the case of surface damage. In order to avoid these situations, case-by-case approaches are necessary.

All plasters, renders or paints that lower the evaporation rate can induce an enlargement in the area affected by the salt solutions. This reduced evaporation rate may be due to a low vapour permeability but also to a low liquid transport capability, as previously discussed. Many industrial plasters and renders present high vapour permeability. But that vapour permeability will hardly surpass that of ancient lime plasters that, moreover, have a higher liquid transport capability. Hence, it is unlikely that salt-accumulating plaster/renders will show a better performance than lime plasters/renders in preventing an increase in the masonry area reached by the salt solutions.

The smaller salt-affected area is obtained with vapour permeable plasters/renders, with high capillary suction (in relation to a given substrate) in which the drying front is, therefore, located as close as possible to the outer surface of the wall. These fast transporting plasters/renders are probably the most adequate solution when protection of adjacent elements is required. However, surface damage is more likely to arise with them.

Only salt-accumulating, salt-blocking or moisture-sealing plasters can provide surfaces free of damage. But for salt-accumulating and salt-blocking plasters, a significant increase in the area attained by the salt solutions may occur. This increase is not necessarily a disadvantage, because the salt load per unit surface is accordingly reduced. Moisture-sealing plasters, on the other hand, can divert the total liquid flux to adjacent areas. Therefore, they are probably only adequate for specific situations such as, for instance, semi-buried walls where the whole surface is to be plastered/rendered. In this case, evaporative crystallization will not take place unless the surface is damaged and escape points arise as represented in Fig. 2.10d.

For salt-blocking plasters, crystallization occurs at the plaster/substrate interface or in the masonry. Therefore, these should be avoided when protection of the masonry is required. Further, they may ultimately induce severe damage patterns such as detachment of the entire plaster/render.

Crystallization on or in the underlying masonry may also arise with plasters and renders without water repellent additives when the suction capability from a certain substrate is not sufficient. As seen in Chapter 4 (Figs. 4.18 and 4.19), with the same plaster L, salt-deposition in the substrate stone arose for the (fine-pored) stone D but not for the (coarse-pored) stone M. The enhanced drying process taking place in this laboratory experiment probably stressed this effect. Indeed, when drying is slower, as it probably happens in buildings, the liquid in the substrate has more time to flow to the plaster. Even so, such occurrence in buildings is probably not impossible, especially in the case of masonry stones with very small capillary pores.

Hence, when protection of the masonry is required, plasters with high suction capability in relation to that substrate should be used. As seen in Chapter 4 (Fig. 4.19), crystallization may also occur in the bedding mortar. Therefore, the plaster should be able to draw, as much as possible, the moisture not only from the (brick or stone) masonry elements but also from the bedding mortars.

In salt-transporting plasters, surface damage can perhaps be reduced when homogeneous pore distribution exists because, in that case, the drying front tends to be sharper. So, if the liquid flux is high enough to maintain the front at the surface, more salt will crystallize as harmless efflorescence rather than in the pores. However, the opposite may happen under conditions such that the drying front is inside the material and, particularly, very close to the surface. In this case, a sharp drying front causes all salt to crystallize just beneath the surface and, thus, detachment of the material surface layer can easily occur.

Slowing down evaporation, for example by the use of a paint layer, could sometimes be positive for salt transporting plasters because surface evaporation and, thus, salt deposition as efflorescence rather than as subflorescence is enhanced.

Plasters/renders may be used, for instance, as provisional sacrificial layers on walls where rising damp was stopped by whichever processes (peripheral drains, damp proof courses, etc.). In this case, a fast transporting plaster/render with strong absorption/drying hysteresis may be appropriate because the drying front will recede later into the plaster with decrease in the total moisture content in the wall. Hence, the masonry will dry faster.

Chapter 6 – Case studies

6.1 Introduction

Salt decay is a serious concern for end users involved in the conservation of monuments and historic urban assemblages. One of the main problems is that salt damage is often worsened after restorations, namely after interventions whose main objective was precisely to mitigate salt decay features. Some possible causes have been put forward in Chapter 5 in the light of the factors that influence the behaviour of plasters and renders in relation to salt crystallization. But a question arises as to whether other factors, more related to practice, could contribute to that worsening in damage. A wrong diagnosis, specific field conditions or application problems could also play a role.

Diagnosis must focus on the two predisposing factors of salt decay: soluble salts and moisture. Yet, as discussed in Chapter 2, preliminary inspections by experts and adequate tests are not normally carried out in many countries, namely, Portugal. It is also possible that, as also pointed in Chapter 2, insufficient background knowledge of the responsible technicians, unclear responsibilities or poor interaction between the involved actors give rise to a wrong approach of the problem, hence, to a poor diagnosis.

Knowledge of the types of salts involved allows their origin to be traced (Goudie and Viles 1997). Valuable information on possible or probable sources of soluble salts can be obtained by study of the enclosure conditions and building history. For instance, in buildings located on the coast there is a high possibility of salt decay due to marine salts deposited directly from the sea, by fog or by contaminated soil water. Also, when a building has been used for storing salty goods it is very likely that salt damage is caused by sodium chloride that penetrated the masonry over time (Nasraoui and Mertz 2004, for example). But not all situations are as straightforward as these and, moreover, different sources and various salts may coexist. And visual observation does not allow distinguishing between different sorts of salts even when efflorescence is visible, which is often not the case.

Moisture sources are also ordinarily identified by visual inspection. Yet, although the experience and scientific background of the expert are a major added value, the physical evidence of dampness is often misleading, as pointed out by Massari and Massari (1993).

In Portugal, traditional plasters and renders are currently used on salt-loaded walls. Pure cement plastering or rendering mortars are becoming less common as it is more and more accepted that they may introduce excessive mechanical stresses or alkali salts in old masonry. Often, a mixture of air-lime and cement is used as binder but, even so, failures occur.

Industrial salt-accumulating plasters and renders are relatively new in Portugal. Hence, there is not enough information from practice to allow assessing their field performance. Yet, due to the practical interest of this working principle and to the high number of (very expensive) products today available on the market, an effort should be made to understand their main possibilities and limitations.

This chapter presents research which was aimed at giving an insight into the field performance of plasters and renders applied on moist salt-loaded walls. Its main objectives were to:

- understand the nature and causes of some serious salt decay problems in old Portuguese buildings, specially as regards the type of salts and sources of moisture;
- identify factors that can account for the worsening of salt decay after restorations;
- compare the possibilities and limitations of diagnostic methodologies based on expertise (namely, visual inspections) or on sampling/testing techniques;
- evaluate the application technique and performance of three industrial plasters for salt-loaded walls available on the Portuguese market.

The research was performed on five old buildings in Portugal: Salvas Chapel (Sines), Alhos-Vedros tide mill (Moita), Despacho House (Pereira), Cloister of Sta. Clara-a-Nova Monastery (Coimbra) and São Sebastião Church (Almada). They were selected, as described in Chapter 2, out of 120 buildings identified by means of a questionnaire sent to 154 Portuguese entities or technicians working in the conservation of old buildings. After preliminary visits to 31 buildings all over the country, the present five cases were chosen. The criteria underlying this selection were both the technical interest, namely intensity of damage and diversity of situations among the five cases, as well as a reasonable proximity to Lisbon which would allow carrying out several inspections at each building.

As with most old Portuguese buildings, these five cases include thick load-bearing ordinary masonry walls. The masonry bedding mortar is, in all cases, weak air-lime mortar. The masonry elements are, in four of the buildings, irregular stone elements of variable size and variable nature. In Despacho House the masonry elements are solid red ceramic bricks.

Diagnosis of the several cases was based on site inspections and on a systematic sampling/testing procedure. HMC and MC profiles were obtained at selected walls of each building, by means of powder drilling, following roughly BRE's protocol (BRE 1989). The salts were investigated by means of X-ray diffraction (XRD) or ion chromatography (IC) analyses.

In São Sebastião Church, test panels of two traditional rendering systems and three industrial systems specific for salt-loaded walls were made on one façade. The performance of the systems was monitored during three years and evaluated by means of visual observation and HMC measurements.

This chapter concerns the research carried out on the five case studies. The buildings and their decay features are described in section 6.2. The diagnosis methodology is explained in section 6.3. The inspections and analyses performed on each building are presented and discussed in section 6.4. The study on experimental test panels is described and discussed in section 6.5. An overall discussion focusing on the objectives of the research is presented in section 6.6.

6.2 Description of the buildings and main decay patterns

6.2.1 - Salvas Chapel

Salvas Chapel (Fig. 6.1) was originally built in the early 16th century, although the front façade dates from the 18th century. It is located in the city of Sines, 150 km south of Lisbon, and is adjacent to Sines harbour (Fig. 6.6). The lateral SE façade (Fig. 6.3) is the one facing the harbour.

At the time of this research, a lime render existed on all façades. According to information from a technician of the national authority responsible for this building, this render included also a (unknown) small percentage of hydraulic cement (unspecified type). This render was applied and lime washed in 1997, in order to replace a previous salt-damaged Portland-cement-based render which dated from around 1986.

The external façades of the chapel showed large decayed spots which reached more than 4m in height from the ground (Figs. 6.1 to 6.4). Damage progressed quickly, as seen in Figs. 6.1 and 6.2.

In October 2003, around 50% of the total rendered area was damaged. The main decay patterns were sanding of the render and further erosion of the render/paint system (Fig. 6.5). No efflorescence was visible. Decayed spots were seen near the ground, but in general degradation occurred mainly at the middle height of the walls (Fig. 6.1 to 6.4). Damage apparently started at the cracks or other singular points of the render (Fig. 6.5).



Fig. 6.1 - Salvas Chapel, SW (front) façade: July 2002 (left), October 2003 (right)



Fig. 6.2 - Salvas Chapel, NE (rear) façade: July 2002 (left), November 2003 (right)



Fig. 6.3 –Salvas Chapel, SE façade, which faces Sines harbour, October 2003



Fig. 6.4 - Salvas Chapel, NW façade October 2003



Fig. 6.5 – Salvas Chapel, damage spots on the SE façade. On the left: cracking of paint/render system and start of erosion, October 2003. On the centre; area in more advanced stage of degradation due to further sanding and crumbling of the paint/render system, October 2003. On the right; detachment of paint and of superficial coat of render, June 2002



Fig. 6.6 - Dona Bataça (water fountain plus reservoir) faces Sines harbour (left) and is located very near to the rear façade of the church (right). The picture on the right is an old photo by DGEMN.

6.2.2 - Alhos-Vedros tide mill

The tide mill (Fig. 6.7) is located in the village of Alhos-Vedros, 10 km south of Lisbon, next to a branch of the Tagus estuary, at less than 10 km from the open sea. The mill is situated between the kettle, an artificial lake meant to accumulate water during the high tide, and the river branch. It dates from the beginning of the 18th century but has been inactive since 1940. It will be rehabilitated to be used as a museum.

In 1999, the existing plasters, apparently based on an (unknown) hydraulic binder, were replaced up to the height of 2.6m on the NW wall, 1.4m on the SW wall and 1.0m on the SE wall, as well as around the NW windows of the ground floor. A traditional hydraulic-lime-based mortar was used, according to information from a technician of the local authority responsible for the building. The whole surface of the walls was then painted with common emulsion paint.

Salt decay was observed mostly in the interior of the mill, on the ground floor. The main decay forms were salt efflorescence, peeling of the paint layer and further sanding of the plaster (Figs. 6.8 and 6.9). Damage affected mainly the new hydraulic-lime plaster. Although infiltration of rainwater was detected on the SE wall (Fig. 6.9), salt damage was more intense and attained greater heights on the NW wall (Fig. 6.8). In general, damage was concentrated close to the ground and around windows on the NW wall. On this wall, damage increased towards the northern corner (Fig. 6.8).

Evidence of dew point condensation was observed on the ground floor of the mill during one of the inspections, as seen and described in Fig. 6.10.



Fig. 6.7 – Tide-mill (N façade)



Fig. 6.8 – Tide-mill, damage at the northern wall of the milling room (ground floor). Left: general view showing that damage increases towards the N corner and appears mostly close to the ground or around the windows. Centre: peeling of the plastic paint around window. Right: damage to paint occurs either without (window frame in the top picture) or with (wall base in the bottom picture) efflorescence.



Fig. 6.9 – Tide-mill, damage on the southern wall. On the left: at the ground floor, salt damage appears mostly on the new hydraulic plaster. On the centre: lack of water tightness of the ground floor windows does not give rise to salt damage. On the right: infiltrations from the roof at the upper floor do not cause salt damage either.



Fig. 6.10 – Tide-mill: moisture stains, probably due to dew point condensation, on the stone pavement of the ground floor. On a dark and rainy autumn day, rain stopped and sun quickly appeared. A much higher temperature was rapidly reached outside the mill, due to the high solar radiation. The temperature inside the mill became clearly lower than outside. The RH inside the mill increased rapidly and moisture spots appeared on the floor.

6.2.3 - Despacho House

Despacho House (Fig. 6.11) is located in the village of Pereira, next to the River Mondego, 200 km north of Lisbon and 20 km from the Atlantic. It was built in the early 18th century, adjacent to Pereira Misericórdia church. Nowadays, it is used as a sacristy and mortuary chapel.

In 2001, the old plasters and renders (unknown type) were replaced by air-lime plasters and renders. It is not clear whether a Portland-cement-based adherence coat was also applied. The walls were painted with a commercial potassium-silicate-based paint that includes water-repellent additives.

Despacho House is located on the flood plain of the Mondego River. It was frequently flooded until the beginning of the 1980s when a system of dikes was finally built to control these periodic floods. The last flood took place in January 2001, precisely during the above-indicated restoration, due to a rupture in the dike system. The flood occurred when the new plaster had already been applied. Despacho House was submerged up to a height of 2.5m but the new plaster suffered no apparent damage. Hence, it was painted in March 2003.

Only a few months after this restoration, degradation in the interior was impressive (Fig. 6.12), affecting around 50% of the plastered area. Damage seems to have the following evolution: (i) efflorescence develops at the interface plaster/paint; (ii) the crystals push the paint layer and eventually cause its rupture; (iii) long needle-like crystals appear on the surface and sanding of the underlying plaster occurs progressively. The plaster shows intense "craquelet" and these cracks are critical points where degradation first starts and develops faster. Damage affects only the upper part of the walls, from a height of 0.7m to 1.5m up to the top of the walls at 2.7m from the pavement.



Fig. 6.11 - Pereira Misericórdia, West façade: tower (left), church (centre), Despacho House (right)



Fig. 6.12 – Extensive damage at the interior of Despacho House. On the left: cracking of the plaster/paint system and efflorescence is observed from around 70cm height from the pavement up to the top of the wall. On the right: damage details showing that salt crystallization at the plaster/paint interface causes rupture of the paint layer and further sanding of the underlying plaster (top), that efflorescence develops preferentially, though not only, on the cracks (centre) and that large areas are eventually covered with efflorescence (bottom).

6.2.4 - Cloister of Sta Clara-a-Nova Monastery

Sta. Clara-a-Nova Monastery is located in the city of Coimbra. It is composed of three independent buildings: monastery, church and cloister. The cloister was built in the first half of the 18th century. It includes the cloister garden and the two-storey surrounding gallery (Fig. 6.13).

The present work concerns the ground floor of the cloister gallery which is a tourist attraction nowadays. Here, a cement-based render, finished with a common white "plastic" paint exists, dating from 1985. This render/paint system shows many salt damaged areas (totalling around 15% of the total rendered area), mostly concentrated close to the ground or located higher on the walls, next to stone elements (Fig. 6.14). Biological growth occurs on the sheltered areas of the walls and vaults (Fig. 6.19).

The damage is mainly sanding of the render and further crumbling of the render/paint system (Figs. 6.15), often preceded by detachment of the paint layer or of a thin superficial render layer which apparently corresponds to the paint penetration depth (Fig. 6.18). Sporadic cracks in the render/paint system are visible (Fig. 6.17) but do not seem to be a general damage enhancement factor. Damage appears to progress according to the sequences in Figs. 6.16 and 6.18.

The roof of the cloister gallery is in poor condition and allows rainwater infiltration. However, no correlation was found between the damaged roof areas and salt damage at the ground floor. The NW wall of the cloister is semi-buried. But salt damage, namely upper spots, also appears on the other walls. The cloister garden has only a partially effective drainage system (Fig. 6.20), which may perhaps lead to rainwater penetration in the upper soil layers beneath the cloister building. Further, probable dew point condensation events were observed during one of the inspections (Figs. 6.19 and 6.20).



Fig. 6.13 - Sta. Clara-a-Nova cloister gallery



Fig. 6.14 – Sta. Clara: damage appears either close to the ground (NE wall, on the left) or higher on the walls, next to stone elements (NW wall, on the right).



Fig. 6.15 – Sta. Clara: details of lower (on the left) and upper (on the centre) damage spots. Efflorescence agglomerations are observed on the upper spot (on the right)



Fig. 6.16 – Sta. Clara: damage development. It is not known whether the borderlines between wet (darker) and dry (lighter) areas and, eventually, between damaged and non-damaged areas correspond to micro-cracks.



Fig. 6.17 – Sta. Clara: sporadic cracks are observed



Fig. 6.18 – Sta. Clara: in some cases, damage starts by the bulging and further detachment of a thin surface layer of the render, accompanied by peeling of the paint, without the presence of efflorescence



Fig. 6.19 – Sta. Clara: moisture spots on the stone (left), moisture spots and biological growth on the render (centre and right, respectively). On February 2004, on a sunny day that followed several cold rainy days, spots of liquid water were observed in the gallery, mainly on the surfaces of the stone columns and arches (left) with further absorption by the render (centre), which suggested the occurrence of dew point condensation. Biological growth is significant on the sheltered parts of the walls and vaults (right), which corroborates this hypothesis.



Fig. 6.20 – Sta. Clara: on the same day in February 2004 probable dew point condensation events were also observed on the stone floor (left and centre). Accumulation of water in the cloister garden (right) had been observed during a previous inspection, in January 2004.

6.2.5 - S. Sebastião Church

St. Sebastian Church (Fig. 6.21) is located in the city of Almada, close to the Tagus River estuary. The present construction was erected in the beginning of the 18th century, over a late 16th century chapel. The church has been damaged and repaired several times throughout its history. It suffered major and minor constructive alterations, went through periods of abandonment and had different functions (it was once used for private housing, as a tavern and even as a stable). In 1999 it was closed for studies and work aimed at its rehabilitation and future reuse as a church.

The church is composed of load bearing ordinary masonry walls around 1.4m thick. The bedding mortar is a very weak and absorbent lime mortar. The stone elements are of a sedimentary nature and include very porous sandstone, calcarenites and argillaceous limestone, from the nearby cliffs of the Tagus River, most of these containing large amounts of fossils of marine life forms.

The church was in very bad general condition at the time of this research. Large areas of the masonry were exposed, due either to the deterioration of the plasters and renders or to their removal. Sanding and crumbling of the masonry bedding mortar (Fig. 6.22) were very severe particularly at the bottom of walls. Erosion depth was up to around 10 cm at several points.



Fig. 6.21 - São Sebastião Church: NE (front) façade in 1999



Fig. 6.22 – São Sebastião: masonry damage progresses as follows: (i) lime plasters and renders disappear due progressive to sanding and crumbling; cement apparently repairs, undamaged, detach; (ii) sanding and crumbling of the bedding mortar progresses on the exposed masonry, as seen in these pictures; (iii) fall of the stone elements eventually leads to masonry disintegration, as seen in the picture on the right.

6.3 Methods

Several inspections of the five buildings were carried out. The information was gathered in a specific form (Annex III), drawn up within the COMPASS project. The sampling / testing methodology used in the several case studies was the following:

- Collection of samples, by powder drilling with 16 or 20 mm rotary drills was done at several heights and depths, in selected walls of each building. Typically, depth indents of 2 cm close to the surface and of 5 cm inside the walls were used, up to a final depth of around 25 to 35 cm.
- The samples were stored in polyetilene bags, carefully sealed to avoid evaporation of the moisture in the samples.
- MC profiles were obtained by measuring the weight loss of these powder samples, after oven drying at 105°C.
- HMC profiles were obtained by exposing these same samples to an environment of 20°C and 95% RH until constant weight was achieved. Narrow-deep receptacles were used here, as mentioned in section 3.6, mostly due to the very large number of sample involved. In this case only a rough and qualitative evaluation of the salt content is possible, as explained in Chapter 3, because the samples are contaminated with not totally known salt mixtures. Yet, here, the HMC values are used mostly to check whether a source of liquid moisture existed in the wall or if the actual moisture in the samples could be solely due to hygroscopicity (BRE 1989).
- IC analyses were performed on some of the most superficial samples (0 to 2 cm or 0 to 5 cm depth). The Na⁺, K⁺, Mg²⁺, Cl⁻, NO₃⁻ and SO₄²⁻ content of these samples was thereby measured.
- Where efflorescence was visible, pure salt samples were collected and subjected to XRD analysis. Also some contaminated mortar samples were subjected to XRD analysis after elimination of the coarse fraction retained in the 106 µm sieve. In this last case, the technique was only expected to help identify the substances that are present in reasonably high amounts (more than around 2% in weight).

The possible water loss during drilling was evaluated by laboratory tests. Five sound bricks from a monument in Lisbon were oven dried at 60°C for 24 hours, weighed and then saturated by total immersion in pure water for one week. Afterwards, they were inserted into polyethylene bags carefully sealed with tape and stored in a conditioned room at 20°C and 65% RH. One month later, the bricks were weighed and then drilled with the same portable drilling device used in the field using a 16 mm rotary drill (Fig. 6.23). Two different drilling velocities were used on different bricks. These velocities were the same as used in the field depending on the hardness of the masonry materials being perforated. As in the field, the process was suspended when excessive heating was detected. The rotary drill was then cooled in water, carefully dried with a cloth and the process resumed. The brick powder was weighed immediately after collection, as well as after a 24-hour oven drying period at 60°C. The initial moisture content of the bricks and that of the drilled powder could therefore be compared. As seen in the graph in Fig. 6.23, no relevant water loss is expected to arise from the drilling process. Indeed, the moisture content in the sampled material is in general even somewhat higher than in the sound bricks. This may be due to a fairly uneven distribution of the moisture in the brick, with a slightly higher concentration of moisture in the core of the bricks where drilling was mostly performed. Although it may be somewhat higher for harder materials, this non-relevant influence of the drilling process on the material moisture content agrees with what is indicated in the literature (BRE 1989, Hall and Hoff 2002).



Fig. 6.23 - Assessment of possible water loss due to the drilling technique

6.4 Results and discussion

6.4.1 - General

In this section the experimental results obtained for each case study are presented. They are then discussed considering also the information gathered during the inspections.

The experimental results include the IC or XRD analysis carried out on selected samples, as well as, for one representative wall in each building (two walls in the case of Sta. Clara) where collection of samples by powder drilling was carried out at several heights and depths:

- a picture of the drilled wall with indication of the sampled heights;
- the type of materials found at the several heights and depths; these materials were evaluated mostly by visual observation of the drilled powder which, then, indicates only their probable nature;
- overall MC/HMC profile (for legibility reasons, only some depths are included);
- in-depth MC/HMC profile corresponding to the lower drilling hole, typically at 0.2m from the ground; for one wall of Sta. Clara an in-depth profile at the height of 3.9 m is also included.

6.4.2 - Salvas Chapel

Collection of samples was carried out in November 2003. The MC/HMC profiles (Fig. 6.24) indicate that rising damp is very significant in Salvas Chapel. The damaging agents are probably chloride and nitrate salts, as seen in Tables 6.2 and 6.3.

Height (m)	Depth (cm)							
	0-2	2-5	5-7.5	7.5-10	10-15	15-20	20-25	25-30
4.0	LM	LM	LM	S	S	LM+S	LM	S+LM
3.0	LM	S	S	S	S	S	S+LM	LM
2.0	-	LM	LM	LM	LM	-	?	LM
1.0	LM	LM	LM	LM	S	S	S	S
0.5	LM	B+LM	B+LM	B+LM	LM+B	LM+B	LM+B	LM+B
0.2	LM	LM	LM	LM	LM	LM	LM	LM

Table 6.1 - Salvas rear façade: sampled materials

LM - lime mortar; S - stone; B - brick

The grey shadows indicate the materials included in the MC/HMC profiles



Fig. 6.24 - Salvas rear façade: drilling location (left) and MC/HMC profiles (centre and right)

Table 6.2 - Salvas rear façade: IC on samples 0-2cm

Low, medium and high anion content (WTA 1991)

Table 6.3 - Salvas rear façade: XDR on fine fraction of render sample collected at 2.1m height and up to 3 mm depth

Halite (NaCl)	+/++
Calcite, CaCO₃	+++
Quartz, SiO ₂	++
Feldspars	vtg/+
Mica	vtg

Fig. 6.24 (centre) shows that the actual moisture content at the base of the wall is very high, much higher than the HMC, and that it decreases with the distance to the ground. This suggests the existence of a source of liquid moisture at the base of the wall. In-depth analysis of the MC/HMC variation in the lower drilling hole (Fig. 6.24, on the right) shows that both the MC and the HMC are constant deep inside the wall, with the MC being much higher (almost 10% higher) than the HMC. These patterns are characteristic of rising damp (BRE 1989, Henriques 1994). It is not known if there is a high groundwater level beneath the church although the nearby presence of a fountain (Fig. 6.6) makes this a likely hypothesis. It is also not impossible that the old reservoir or old fountain pipes are damaged and leak considerable amounts of water.

The damaging salts deposit preferentially close to the maximum level(s) attained by the moisture in a wall (Arnold 1982). The present MC/HMC profiles show that, in fact, the stronger accumulation of salt occurs at around 2m from the ground.

IC (Table 6.2) indicates that chloride and nitrate (probably of sodium) are likely to be the responsible salts. XRD analysis of a render sample collected at the height of maximum salt content indicated halite as the main salt (Table 6.3). The nitrate is likely to originate from the soil. The chloride may have the same origin (the groundwater may be contaminated by sea water), be deposited by salt mist and further absorbed, either into the soil or directly by the wall.

6.4.3 - Alhos-Vedros tide mill

Collection of samples was carried out in November 2003. In this case and despite the direct contact of the mill foundations with the river water (Fig. 6.7), rising damp does not seem to be the fundamental problem here (Fig. 6.25). The damaging salts are mostly sodium carbonate and bicarbonate salts, together with some sodium chloride (Table 6.5).

Height (m)	Depth (cm)							
neight (m)	0-2	2-5	5-7.5	7.5-10	10-15	15-20		
2.7	HM	HM	LM+HM	LM	LM	LM		
2.0	HM	HM	LM+HM	LM	LM	LM		
1.5	HM	HM	LM+HM	LM	LM	В		
1.0	HM	HM	S	S	S	S		
0.5	HM	HM	LM+S	LM+S	LM+S	LM+S		
0.15	HM	HM	LM+HM	LM	LM+S	LM+S		

Table 6.4 - Tide mill NW wall: sampled materials.

HM – hydraulic mortar, LM – lime mortar; S – stone; B – brick



Fig. 6.25- Tide mill NW wall: drilling location (left) and MC/HMC profiles (centre and right)

Termonatrite - Na ₂ CO ₃ .H ₂ O	+
Gaylussite - Na ₂ Ca(CO ₃) ₂ .5H ₂ O	+
Trona - Na ₂ H(CO ₃) ₂ .2H ₂ O	+
Halite - NaCl	+
Calcium hydroxide - Ca(OH) ₂	+
Calcite - CaCO ₃	+
Quartz - SiO ₂	++
Feldspars	+
Міса	+
Caulinite	+

Table 6.5 - Tide mill NW wall: XDR on efflorescence collected up to 0.5m in height from the pavement on the drilled area

Both the overall MC/HMC profiles and the in-depth MC/HMC profile at 0.15m height (Fig. 6.25) show that, deep inside the wall, the MC is low and similar to the HMC at 95% RH. Therefore, rising damp in the wall is probably low, part of the actual moisture being of hygroscopic origin. Although detailed information on the mill foundations is not available, visual observation from the exterior (Fig. 6.7) suggests that large stone blocks were used with little or no significant presence of bedding mortars. Therefore, it is likely that the rise of damp is drastically hindered by these open joints.

In Alhos-Vedros tide mill, decay is mostly confined to the new plaster (Figs. 6.8 and 6.9). XRD analysis (Table 6.5) showed that, although sodium chloride is present (probably carried by some capillary rising moisture or by salt mist), efflorescence is mainly composed of alkalicarbonate salts. Alkali-carbonate salts are possibly derived from the hydraulic-lime used in the plaster (Arnold 1982, Arnold and Zehnder 1989). This new plaster was used to repair the bottom areas of the walls and those around the NW windows of the ground floor. The fact that the repair involved much larger areas on the NW than on the SE wall suggests that salt damage in the mill was also more intense at N before this restoration. The higher damage at the NW wall, around the windows and, particularly, at the North facing corner, suggests a relation between the occurrence/evolution of damage and the fact that these are colder areas, where evaporation is lower and the risk of dew point condensation is higher. The hypothesis of dew point condensation being a key source of moisture (Gonçalves *et al* 2005) in the mill was corroborated by direct observation during one of the inspections, as seen and described in Fig. 6.10.

Dew point condensation may occur on or in cold walls. Particularly when the air is humid, namely during or after rainy days, if the outside air temperature rises due to solar radiation, the absolute humidity increases because hotter air can hold a higher amount of water vapour. When this air comes in contact with colder surfaces its temperature drops. In consequence, the relative humidity of the air increases. If the saturation humidity is exceeded, dew point condensation occurs.

Posterior research (Cardoso 2006), to which the author contributed, was performed on a similar tide mill, Corroios tide mill, also located in the Tagus estuary. This research included environmental monitoring over a period of 14 months by means of a thermohygrometer. Very high RH was systematically measured, namely in the milling room which is similar to that of Alhos-Vedros tide mill. The maximum RH in this room was between 75% in July and more than 90% in November and December. Although the time scale was probably not detailed enough for sporadic condensation events to be detected, it is likely that such a high RH in the middle of the room could give rise to surface condensation on the walls that were probably often colder than the surrounding air.

6.4.4 - Despacho House

Collection of samples in Despacho House was carried out in April 2004. As in Alhos-Vedros tide mill, the salts are mostly alkali (sodium) carbonate salts (Table 6.7). But here, differently, the moisture content in the walls is rather high (Fig. 6.26).

Indeed, both the overall MC/HMC profiles (Fig. 6.26, centre) and, particularly, the in-depth MC/HMC profile at 0.2m height from the pavement (Fig. 6.26, right) show that, deep inside the wall, the MC is more than 7% higher than the HMC at 95% RH.

Height (m)	Depth (cm)							
	0-2	2-5	5-10	10-15	15-20	20-25	25-30	
2.6	M1	B+LM	В	B+LM	LM+B	LM+B	LM+B	
2.1	M1	B+LM	B+LM	B+LM	B+LM	LM	LM	
1.5	M1	LM	LM+B	LM+B	-	LM	LM	
0.7	M1	LM	LM	LM	LM+B	LM	LM	
0.2	M1	LM	LM	LM	LM	LM	LM	

 Table 6.6 - Despacho internal wall: Sampled materials

M1 – (lime?) mortar, LM – lime mortar; B - brick



Fig. 6.26 - Despacho internal wall: drilling location (left) and MC/HMC profiles (centre and right).

Sampling he		ı height (r	eight (m)		
1.1 1.5 2.1 +++ + - + +++ - - + +	2.1	2.6			
++	+	-	-		
+	++	-	-		
-	+	+	++		
-	-	vtg	-		
?	?	+	+		
+	+	++	+/++		
+	+	++	++		
	1.1 ++ - - ? + +	Sampling 1.1 1.5 ++ + + ++ - + - - ? ? + + + +	Sampling height (n 1.1 1.5 2.1 ++ + - + ++ - - + + - - Vtg ? ? + + + ++ + + ++		

Table 6.7 - Despacho internal wall: XDR on efflorescence collected on the same internal wall

It is not clear if this is a normal case of rising damp or if there was also a relevant accumulation of water inside the walls during the 2001 flood. The latter is suggested by the in-depth profile (Fig. 6.26, on the right) which shows that the MC clearly increases towards the surface, nonetheless the HMC is overall constant from the 10-15 cm depth up to the surface. In any case, the water repellent paint is possibly lowering the wall drying rate and, therefore, inducing larger damaged areas: water repellent layers only permit vapour migration and, hence, "push" the drying front into the material.

XRD shows (Table 6.7) that efflorescence is mainly composed of alkali (sodium)-carbonate salts. Hence, as in Alhos-Vedros tide mill, the main source of salts is probably the plaster/paint system, though it was not possible to know whether it is the plaster or the silicate-based paint.

Similarly to Salvas Church where very high moisture content also exists in the masonry, damage is concentrated on the upper and not on the lower part of the wall. In the case of Salvas Church the salts originate (at least partially) from the ground. It is probable that the ground solutions only reach saturation after considerable evaporation, that is, further on their way up the wall. Here, in Despacho House, it is likely that the high moisture content at the base of the walls is preventing crystallization of the carbonate salts.

6.4.5 - Cloister of Sta Clara-a-Nova Monastery

Collection of samples was carried out in November 2003 (Fig. 6.27) and April 2004 (Fig. 6.28). On-site observations showed that damage systematically starts next to stone elements (window frames, arches or pavement), as seen in Fig. 6.14, and that Sta. Clara stones "exude" liquid water after cold days, when the air heats up (Figs. 6.19 and 6.20). Dew point condensation on the stone elements or hygroscopic absorption by these elements were therefore considered possible sources of moisture.

In the case of the upper damage spots, infiltration from the first floor was also a possibility, since both the floor and roof on the first floor were in bad condition. For the lower damage spots, infiltration of rainwater from the cloister garden, which has deficient drainage conditions, into the upper soil layer was another hypothesis.

Height (m)				De	pth (cm)			
	0-2	2-5	5-10	10-15	15-20	20-25	25-30	30-35
4.9	CM	CM+LM	LM	LM	LM	LM	LM	LM
3.9	СМ	CM	S	S	S	S	S	S
3.4	CM	CM+LM	LM	LM	LM	LM	LM	LM
2.4	CM	CM	S	LM	-	LM	LM	LM
1.4	CM	CM+S	S	S	S	_	LM	LM
0.5	CM	CM	S+CM	S	S	S	S	S

Table 6.8 – 3	Sta, Clara	NW wall:	Sampled	materials
10010 0.0	olu. olulu	TTTT TOTT	Cumpicu	matorialo

CM – cement mortar; LM – lime mortar; S – stone



Fig. 6.27 - Sta Clara NW wall: drilling location (left) and MC/HMC profiles (centre and right)

In the case of the upper damage spot (Fig. 6.27), the overall MC/HMC profile and the in-depth profile at 0.2m height from the ground suggest that rising damp is very limited in this wall. As seen in the in-depth profile, the actual moisture content deep inside the wall is only

around 2% higher than the HMC. Therefore, the occurrence of damage at such high distances from the ground is hardly assignable to rising damp. Further, since this wall was drilled during the rainy season, it is not likely (though certainly not impossible) that much higher moisture contents exist in other periods.

The in-depth MC/HMC profile (Fig. 6.27, right) at the height of maximum damage (3.9m) shows that moisture decreases towards the interior of the wall and that, at the deepest points, the MC is similar to the HMC. A superficial source of liquid moisture is therefore likely to exist.

Height (m)	Depth (cm)							
ricigiit (iii)	0-2	2-5	5-10	10-15	15-20	20-25	25-30	
4.0	CM	LM	RM	RM	RM	RM	RM	
2.5	CM	LM+S	S	S	S	S	S	
1.5	CM	LM+S	S	S	S	S	S	
0.7	CM+S	S	S	S+LM	S+LM	S+LM	S	
0.2	CM	LM+S	S+RM	RM	RM	RM	RM	

CM - cement mortar; LM - lime mortar;

RM – coarse argilaceous mortar; S – stone; B – brick



Fig. 6.28 - Sta Clara NE wall: drilling location (left) and MC/HMC profiles (centre and right)

In the case of the wall with damage close to the pavement (Fig. 6.28, centre), the overall MC/HMC profile shows that the MC decreases with height. However, the in-depth MC/HMC profile (Fig. 6.28, right) shows that the actual moisture content (MC) decreases inside the wall towards its interior and that, at the deepest point, the MC is lower than the HMC. A superficial source of liquid moisture is therefore likely to exist also in this case.

The similarity of the in-depth profiles of the two walls at the height of maximum damage is interesting because they concern completely different heights: 3.9m (Fig. 6.27) or 0.2m (Fig. 6.27). Absorption of (hygroscopic or condensed) moisture from the stone elements

(upper damage spots) or from the ground (lower spots) seems, thus, a strong hypothesis. However, is possible that, for example, the water used for washing the stone pavement has also a contribution in the case of the lower spots.

Dew point condensation can occur when hotter air comes into contact with colder walls because the air RH increases and the saturation humidity may thereby be exceeded. This mechanism can also account for hygroscopic absorption by salt contaminated walls. Indeed when the air RH increases next to colder surfaces, the RH_{eq} of the contaminant solutions can be surpassed and hygroscopic absorption may therefore take place.

XRD (Table 6.10) indicates a strong presence of alkali-carbonate salts both in the upper and in the lower damaged spot. In the lower spot some nitrate was also identified. It is likely that the carbonate salts originate from the cement render itself. The nitrate probably comes from the soil. This suggests that a (low) influx of moisture from the ground exists which is in line with the slight decrease of the MC over height (Fig. 6.27 and 6.28) possibly due to bad drainage of the cloister garden or to penetration of water during cleaning of the pavement.

High nitrate contents were also found in the lower spot by IC (Table 6.11). In the upper damage spot the IC revealed somewhat lower nitrate content. The significance and origin of the nitrate in this upper damage spot are unknown.

	NW wall (upper spot)	NE wall (lower spot)	
Crystalline compounds	Efflorescence 1.5 m from ground	Sanded surface material 0.2 m from ground	
Trona, Na ₃ H(CO3) ₂ .2H ₂ O	++/+++	?/vtg	
Gaylussite, Na ₂ Ca(CO ₃) ₂ .5H ₂ O	-	+	
Niter,KNO ₃	-	?/vtg	
Calcite, CaCO ₃	++/+++	+++	
Dolomite, CaMg(CO ₃) ₂	-	+	
Quartz, SiO ₂	vtg	+	

Table 6.10 – Sta.	Clara: XDR or	n efflorescence or	sanded	surface materia
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able 6.11 – Sta. (Clara: IC on	samples 0-	-2cm
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Façade	Height (m)	Ion content (% W)					
		Na⁺	K⁺	Mg ²⁺	Cl	NO ₃ ⁻	SO42-
NE (lower spot)	2.5	0.05	0.11	not detected	0,03	0,02	0,07
	0.7	0.31	0.26	0.04	0,32	0,71	0,16
	0.2	0.35	0.25	not detected	0,43	1,71	0,17
SE (upper spot)	4.2	0.72	0.60	not detected	0,37	0,22	0,11
	3.0	0.38	0.25	not detected	0,46	0,42	0,18
	0.2	0.47	1.27	not detected	0,10	0,16	1,47

Low, medium and high anion content (WTA 1991)

6.4.6 - S. Sebastião Church

Sampling was carried out in November 2003. The walls were found to be practically dry (Fig. 6.29) and the salts seem to be mostly nitrate salts (Table 6.12).

Height (m)	Depth (cm)						
	0-5	5-10	10-15	15-20	20-25	25-30	30-35
3.5	LM	LM	LM+S	LM+S	S	S+LM	S+LM
2.5	LM	LM+S	LM+S	LM	LM	LM	S
1.5	LM	LM	LM+S	S	S	S	S
0.5	LM	LM+S	S	S	S	S	S
0.2	LM	LM	LM	LM	LM+S	LM+S	LM+S

Table 6.12 - S. Sebastião exterior: sampled materials

LM – lime mortar; S – stone



Fig. 6.29 - S. Sebastião: drilling location (left) and MC/HMC profiles (centre and right)

Height (m)	Ion content (% W)						
	Na⁺	K⁺	Mg ²⁺	Cl	NO ₃ ⁻	SO4 ²⁻	
1.5	0.06	0.01	nd	0.12	0.12	0.06	
0.5	0.27	0.13	nd	0.48	0.71	0.49	
0.2	0.34	0.19	nd	0.54	1.01	0.26	

Table 6.13 – S. Sebastião exterior: IC on samples 0-2cm

Low, medium and high anion content (WTA 1991)
As seen in the MC/HMC profiles (Fig. 6.29), the HMC and the MC profiles have similar shapes, but the HMC values are clearly higher. This indicates that the actual moisture content in the wall is probably of hygroscopic origin. The decrease in the HMC with height indicates that the salts come from the ground, although the rising damp mechanism was probably not active at the time of sampling. The salts concentrate close to the ground, as indicated by the decrease in the HMC with height, which suggests that the moisture contents involved in salt transport and deposition are not, at least permanently, very high.

Nitrate seems to be the main ion involved (Table 6.13), which is consistent with the hypothesis of capillary rise from the ground. It is interesting that, despite vestiges of marine life forms exist in the masonry stone, this masonry does not seem to be a relevant source of salts.

6.5 Research on experimental test panels

6.5.1 - Materials and methods

Test panels of two traditional rendering systems and three industrial systems specific for salt-loaded walls were made (Fig. 6.30) on the NW façade of S. Sebastião Church.



Fig. 6.30 – Test panels: traditional renders A and C and industrial renders D, E and F

A and C are traditional rendering systems based on air-lime or air-lime and cement, respectively. D, E and F are industrial systems currently commercialised in Portugal that are claimed to be "dehumidifying" systems, that is, systems able to provide a final surface free of moisture and salts. The stated working principle of systems D and E is salt accumulation. The working principle of system F is not clear but its (stated) water repellent properties suggest that it is probably a salt-blocking system.

The test panels had flat dimensions of around 2m x 2m. Their characteristics are described in Table 6.14.

Panel (system)	Coat application order	Mortar	Volumetric composition / expressed working principle	Coat thickness
A	1 st	А		15 mm
(traditional air-lime render)	2 nd	А	1 : 1.5 : 1.5 (Lime : ST : SC)	15 mm
С	1 st	C1	1 : 1 : 6 (CII : Lime : ST)	5mm max*
(traditional	2 nd	C2	$1 \cdot 2 \cdot 12$ (CII · Lima · ST)	15 mm
render)	3 rd	C2	1.3.12 (CII.LIME.ST)	15 mm
D	1 st	D1	Adherence coat Calcareous aggregates, hydraulic binders without C3A and additives (for promoting adherence and for obtaining a high porosity and hydro-repellence)	10 mm
(industrial salt-accumulating system)	2 nd	D2	Accumulating coat Based on white natural hydraulic lime and lightweight	25 mm
	3 rd	D2	aggregates (hydrated silicates of expanded aluminium)	25 mm
	4 th	D3	Finishing coat Fine white mortar based on natural hydraulic binders, air lime and sand (does not include cement)	10 mm
E	1 st	E1	Adherence coat Hydraulic lime, pozzolana, siliceous and calcareous aggregates	5 mm
(industrial salt-accumulating system)	2 nd	E2	<u>Accumulating coat</u> Hydraulic lime, pozzolana, siliceous and calcareous aggregates	15 mm
	3 rd	E3	Finishing hydro-repellent coat (grey) Hydraulic lime, pozzolana, siliceous and calcareous aggregates	5 mm
F (industrial system	1 st	F1	<u>Adherence coat</u> Hydraulic lime, siliceous sand	5 mm
with no clear	2 nd	F2	Hydro-repellent coat (grey)	20 mm
working principle)	3 rd	F2	Hydraulic lime, siliceous sand	20 mm

Table 6.14 - Rendering s	systems in th	e test panels
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Lime – Dry hydrated lime LUSICAL; CII – Portland cement (type II); ST – Sand from the Tagus River SC – Yellow pit sand from Corroios; * Discontinuous coat

The mortars that composed the three industrial systems, as well as the air-lime mortar of system A which served as a reference, were characterized by means of capillary unidirectional absorption tests (CEN 2002). These results are presented in Fig. 6.31. D3 mortar was not tested, in the first place, because there was not enough product available and, ultimately, because its capillarity coefficient was not necessary for interpreting the behaviour of the systems, as it will be seen in section 6.5.3.



Fig. 6.31 – Capillary water absorption of the three industrial systems and traditional mortar A. The numbers in brackets correspond to the capillarity coefficient, in kg.m⁻².h^{-1/2}.

It is interesting to note that, for mortars D2 and E2 which correspond to the layer where salts are supposed to accumulate in either system, the absorption curves show several distinct slopes (Fig. 6.31). The same had been observed for salt-accumulating MEP and Parlumière plasters (Fig. 5.5). These multi-slope absorption curves indicate the existence of distinct migration kinetics in the same porous material, as explained in section 4.2.2 and depicted in Fig. 4.3. Also water-repellent mortars E3 and F2 show different absorption slopes, albeit less markedly. On the contrary, mortars D1, E1 and F1, similarly to traditional lime mortar A, only show an initial dominant slope.

Performance of the rendering systems was monitored during three years, by visual observation of the tests panels. At the end of this period, HMC measurements were carried out:

- on bedding-mortar or stone pieces collected at the surface of the masonry, to allow comparing the salt-content of the two types of material;
- on samples collected by powder-drilling across each rendering system, in order to obtain in-dept HMC profiles that could disclose the salt-accumulating behaviour of the systems.

Collection of samples by powder drilling and the HMC measurements were carried out as described in section 6.3.

6.5.2 - Execution of the test panels

Before making the test panels, the masonry was cleaned, as illustrated and explained in Fig. 6.30. These cleaning operations together with the application of the 15 layers that compose the five rendering systems (Table 6.14) were carried out over a total of 10 working days and over a period of two months.



Fig. 6.32 – Before making the panels, the substrate was prepared as follows: (i) the old lime render, in poor condition and with many lacunas, was removed; (ii) the old lime mortar in the joints between masonry stones was removed up to a depth of around 15mm; (iii) the masonry was brushed to eliminate as much loose material as possible; (iv) all recesses deeper than 30 mm were filled with lime mortar A and small stones of the original masonry stone type. All systems, except industrial system E, were applied by a mason with general experience in plastering and rendering. The application of industrial systems D and F was carried out under direct instructions of technicians from the supplying companies. Industrial system E was mostly applied by a specialized mason who usually does practical demonstrations with this particular product. The general mason collaborated with him and was able to learn the technique through practice. After some time during the application of each coat, the general mason was able to carry out the application on his own.

The application of both the traditional rendering systems and industrial system E were successful. Only in the case of mortar E2 was the recommended amount of mixing water found to be too high but the problem was promptly detected and corrected by the specialized mason. In contrast, numerous problems also arose during the application of industrial systems D and F:

- In some cases (D1 and D2), the recommended amount of mixing water was found to be insufficient for achieving an acceptable workability. New mixings had to be done and, for this reason, the available product was only sufficient for a 1.5 m wide panel.
- One mortar (D2) was initially applied in a thickness higher than the maximum recommended in the written documentation (30 mm). The fresh layer detached (Fig. 6.33 iv) and fell down. The application had to be repeated by using two coats of 25 mm applied with a one day interval.
- Mortar D3 cracked immediately after application (Fig. 6.33 vi).
- It was very difficult to finish the last layer of mortar F2 (smoothening with trowel). The fresh mortar stuck to the trowel, even two hours after the application, and the operation systematically led to a very irregular surface (Fig. 6.33 ii). The next day, more mortar was prepared to try to fill in the numerous cavities on the surface. But the operation was not fully successful and even after the completion of this procedure the final surface was much more irregular than reasonable or expected.



Fig. 6.33 – Test panel execution: (i) application of the last coat of traditional lime render A; (ii) irregular surface of industrial F2 layer; (iii) application of industrial E3 layer; (iv) detachment of fresh D2 layer; (v) application of industrial D3 layer; (vi) crack on fresh D3 layer.

6.5.3 - Evaluation of the systems after three years of natural exposure

Three years after the applications, there was apparently no damage by salt crystallization in any of the test panels. However, cracking was observed on all systems except lime-cement render C (Fig. 6.34):

- A and E showed some disperse and long cracks apparently caused by differential movement of the masonry;
- F showed numerous micro-cracks;
- D3 showed intense "craquelet", as well as very intense accumulation of dirt.



Fig. 6.34 - Cracking of the rendering systems three years after their application

The deposition of salt across the systems was evaluated by means of HMC profiles. Sampling was performed in such a way as to always obtain the bedding mortar as substrate material because salt deposition seems to be more significant in this bedding-mortar than in the masonry stone (Fig. 6.35). Nonetheless, the HMC profiles (Fig. 6.36) show that the salt content in the substrate varied significantly from panel to panel. Hence, only a rough comparison of the different systems is possible.

These HMC profiles (Fig. 6.36) indicate that:

- The three industrial renders (D, E and F) and also the traditional lime-cement render (C) seem to be able to prevent salts from reaching the surface. In contrast, the traditional pure lime render (A) transported the salts to the surface.
- Lime-cement render C tends to accumulate the salts in the adherence coat.
- The systems with a very absorbing first coat (A and D) seem to be more able to transport the salts from the substrate to the render: in both cases, the substrate and the first render coat have similar HMC values.
- The systems with a low absorbing first coat (E and F) seem to induce a more significant deposition of salt in the substrate: the HMC decreases from the substrate to the first render coat.





Fig. 6.36 - In-depth HMC profiles at 0.4m height from the ground

6.6 Discussion

6.6.1 - Main causes of decay

Table 6.15 summarizes the decay patterns and main diagnostic hypothesis for the five case studies.

Case study	Type of plaster/render		Salts		Probable origins
of the decay		Main degradation patterns	Nature	Probable origins	of moisture
Salvas Chapel <i>Exterior</i>	Traditional lime render (with some cement) and lime wash	 Cracking of render/paint system (damage develops preferentially in the cracks) Sanding and erosion of the render/paint system Most decayed spots are located at the middle height of the walls. 	 Halite Nitrate (probably of sodium) 	 From the soil The chloride may also be deposited by salt mist 	Rising damp
Alhos- Vedros tide mill Interior	 Efflorescence, peeling of paint and sanding of plaster Damage close to the pavement and around the NW 	Alkali (sodium) carbonate salts	Hydraulic plaster	Dew point condensation and some rising damp	
	windows - Damage more intense at N	Halite	River water		
Despacho House <i>Interior</i>	Traditional air-lime plaster and potassium silicate-based paint with water repellent additives	 Cracking of render (damage develops preferentially in the cracks) Efflorescence, blistering and rupture of paint, sanding of render Degradation appears on the upper part of the walls 	Alkali (sodium) - carbonate salts	Plaster / paint system	Rising damp (accumulation of water during the 2001 flood is also possible)
Sta. Clara cloister <i>Exterior</i> Traditional cement-based render and common emulsion paint	 Detachment of the paint or of a render layer corresponding to the paint penetration depth Sanding of render, crumbling 	Alkali carbonate salts	Render	- Condensation or hygroscopic absorption by the stone and further migration to the	
	 of render/paint system Degradation close to the ground or higher on walls, next to stone elements 	Nitrate and chloride	Possibly the soil	render - Penetration of ground water due to bad drainage of the garden	
S. Sebastião Church Interior and exterior	- Traditional lime plasters and renders. - Cement- based repairs	 Sanding and crumbling of lime plaster and renders Detachment of cement repairs Sanding and crumbling of the masonry bedding-mortar 	Chloride and nitrate	From the soil	Minor rising damp

Table 6.15 - Summary of decay and diagnosis

Four of the five case studies (Salvas, Alhos-Vedros, Despacho and Santa Clara) show pathological decay situations with the following likely causes:

- Crystallization of alkali-carbonate salts was the fundamental cause of decay in three out of the five case studies: Alhos-Vedros, Despacho house and Santa Clara. As explained in section 2.2.4, carbonate salts originating from high alkali content materials are often involved in salt decay processes (Schaffer 1932, Arnold 1981). It is likely that the use of high alkali content materials is a recurrent cause of salt decay in Portugal. This type of decay arises probably in many cases from the use of cement-based mortars. However, in the case of Despacho House it is not know whether the silicate-paint had any contribution. Two of the case studies (Alhos-Vedros and Despacho House) concerned recent restoration interventions. This suggests that the use of high alkali content materials can probably account for several of the cases where damage is worsened after restoration.
- Significant rising damp was found in Salvas Church and Despacho House.
- Surface dampness was apparently the main source of moisture in Alhos-Vedros and Santa Clara. The moisture arises probably from dew point condensation, as explained in sections 6.4.3 and 6.4.5. However, as pointed in section 6.4.5, hygroscopicity is another possible source of moisture. In both monuments, the damaging salts were alkali carbonate salts. It is possible that surface moisture is a particularly effective decay agent in cases such as these where the salts form mostly at the exposed surface of the walls.
- Cracking of plastering or rendering systems seems to worsen surface decay. This was observed in Despacho House and Salvas Church: the cracks seem to be critical points where damage preferentially starts and progresses faster. Indeed, although large cracks are mostly expected to enhance evaporation, capillary cracks may enhance liquid transport to the surface and, hence, locally accelerate the decay rate. In both buildings, crack networks (the so-called "craquelet" pattern) are observed (Figs. 6.5 and 6.12). This pattern is typical of plasters and renders with a strong drying shrinkage and is usually attributed to an excess of binder or of fine aggregates in the mortar, application of too thick plaster/render layers, too dry substrate, application carried out during hot and dry weather or direct incidence of solar radiation on the fresh mortar (Veiga 2004, for instance). It is not known whether in Salvas Church and Despacho House any of these factors was significant. But one common feature of these two buildings is the existence of very high moisture contents in the masonry. Whether this factor could contribute to enhance the drying shrinkage, perhaps somewhat similarly to an excess of mixing water, is a hypothesis that requires further research.

S. Sebastião Church is a different case. There is no evidence of this being a pathologic situation as regards salt decay even though reasonable amounts of nitrate and chloride were found at the base of the wall. These salts are likely to come from the soil. Nitrate, in particular, may be related with the church having worked as a stable, as mentioned above. This masonry is probably subjected to seasonal events of capillary rise of moisture from the ground but nothing points at high amounts of moisture being involved. All things considered, this is probably a normal situation for an old building with thick masonry walls built in direct contact with the ground and not far from the sea. Damage seems to result mostly from the absence of adequate (sacrificial) plasters and renders and, ultimately, from a total lack of maintenance.

6.6.2 - Diagnosis

The work presented in this chapter demonstrates that an adequate diagnostic methodology includes both visual inspection and sampling/testing.

Indeed, "blind" sampling/testing will generally not be enough namely because sophisticated techniques, for instance of environmental monitoring, are normally not available nor are they feasible. Therefore, expertise can provide valuable complementary information on the causes of damage.

For instance, in Sta. Clara and Alhos-Vedros the possible sources of moisture were identified by observation and further interpretation of the visible damage patterns.

However, simple visual observation may sometimes lead to wrong conclusions. For example, in Alhos-Vedros tide mill, whose foundations are in direct contact with water, rising damp would be a straightforward hypothesis as regards the source of moisture. But the MC/HMC profiles indicated that rising damp is limited and, hence, a different source of moisture had to be sought. Indeed, as stated by Massari and Massari (1993), as far as dampness is concerned, "reality often contradicts good sense and the obvious hypotheses are frequently false". Moreover, it is not possible to distinguish visually the type of salt involved in a decay process. And, as seen in the cases of Alhos-Vedros, Despacho and Santa Clara, an accurate diagnosis is often only possible when that information is available.

Diagnosis must focus on the two predisposing factors of salt decay: soluble salts and moisture. However, in Portugal as in other European countries (Gonçalves 2002b, TNO 2002), it is not common to perform preliminary inspections based on an oriented methodology and with adequate auxiliary methods of diagnosis. Hence, it is likely that salt and moisture sources are often not properly identified. For example, in Salvas Church and Despacho House, no adequate measures to prevent rising damp were adopted in the recent restorations because the situation was not properly diagnosed. Further, in none of the cases where carbonate salts were present, the hypothesis of the plastering or rendering system being the main source of salt had ever been put because the type of salt had not been determined by adequate analyses.

6.6.3 - Test panels

Although the use of test panels may allow a more realistic evaluation of the behaviour of plasters and renders in relation to salt crystallization, the work done in S. Sebastião Church shows that these are not straightforward tests: (i) it takes considerable time and effort to properly prepare the substrate and make the test panels; (ii) years of exposure may be necessary to obtain visible damage; (iii) moisture and salts are often heterogeneously distributed in walls and this may complicate the analysis of the results .

Application of the industrial renders made it clear that, although well written or even oral instructions are certainly important, they are not enough to ensure successful application. Plasterers must also gain practical experience with the products. Indeed, it was not possible to correctly apply two of the systems (D and F), even when the application was done by an experienced mason under direct supervision of technicians from either supplying company. The third system (E) was applied by a mason from the supplying company. This mason had specific training and usually does practical demonstrations with his product. The application was able to learn the technique from his colleague.

Chapter 7 – Summary, conclusions, and future perspectives

7.1 Summary

This thesis was aimed at understanding the behaviour of plasters and renders on salt-loaded walls. The research was carried out through the following main steps:

- The current state-of-the-art and state-of-the-practice in salt decay features were accessed focusing particularly on old plastered/rendered buildings and their conservation practice in Portugal.
- Two general laboratory techniques, for RH control and salt content measurement, were investigated in order to decide whether and under which conditions they could be adequate experimental tools.
- Drying of salt-loaded materials was studied which allowed a better understanding of the influence of soluble salts on the drying of porous materials, mechanisms of salt-induced dampness and evaporation-induced salt distribution in masonries.
- The behaviour of plasters and renders in relation to salt crystallization was, afterwards, investigated. By this work, the influence of various factors (type of salt, plaster properties, nature of the substrate material, presence of a paint layer, use of water-repellent additives or the material saturation condition) was assessed. Hypotheses were presented to explain why sodium chloride is typically much less damaging than sodium sulfate in laboratory tests. It was also possible to identify several factors that can account for the worsening of salt decay problems after restoration interventions. Further, the possibilities and limitations of salt crystallization tests as performance tests for plasters and renders were discussed. Also, general guidelines for the choice of appropriate plasters and renders for salt loaded walls were presented
- Finally, the study of five old buildings in Portugal provided an insight into the field performance of plasters and renders applied on moist salt-loaded walls. It was possible to understand the nature and probable causes of decay, to compare expertise-based and sampling/testing-based diagnostic methodologies, as well as to evaluate the application technique and performance of different rendering systems applied as test panels in S. Sebastião church. As these case studies are expected to represent many old buildings in Portugal, further hypotheses could be made on factors that may account for the worsening of salt damage after restoration interventions.

This research was based on the tasks described in the next paragraphs.

A questionnaire asking for information on salt decayed plastered/rendered buildings was sent to 154 entities and professionals in Portugal. Through this questionnaire and some further personal contacts, 14 end-users were selected to be interviewed and 31 buildings out of the 120 indicated were identified as possible case studies.

These 14 end-users were interviewed (Gonçalves 2002b, Gonçalves *et al* 2003) to provide an insight into how salt decay problems are approached in practice, as well as into the types of plasters and renders used on salt-loaded walls.

A survey was carried out in the Portuguese market (Gonçalves 2002a) to identify industrial plasters/renders or related products available for moist salt-loaded walls. This survey also allowed selecting three industrial systems to use in the test panels of S. Sebastião church.

Salt decay pathology was structured and discussed. Salt decay anomalies, signs, physical medium, causes, mechanisms, predisposing and adjuvant factors were addressed mostly by means of a review of state-of-the-art literature. The discussion addressed also the main working principles of plasters and renders for salt-loaded walls, as well as the subject of salt crystallization tests.

An overview of the state-of-the-practice on handling of salt decay problems in plastered/rendered buildings was given. It was largely based on the above mentioned interviews and market survey and focused mainly on the case of Portugal.

Background information was presented on specific topics such as hygroscopicity, moisture transport and drying of porous materials, main variables in salt crystallization tests and sorption behaviour of porous building materials.

The possibilities and limitations of using aqueous solutions to generate a constant RH with salt-loaded materials were assessed by means of experimental research on NaCl (Gonçalves and Abreu 2007).

The possibilities and limitations of the HMC method for salt content evaluation were investigated, both on a theoretical basis (Gonçalves and Rodrigues 2006) and by means of experimental research on nine different salts (Gonçalves *et al.* 2006a).

Drying of salt-loaded porous materials was investigated (Gonçalves *et al.* 2006c) mostly by means of MRI-monitored drying experiments performed on specimens composed of stone, bedding mortar and plaster.

The behaviour of plasters and renders in relation to salt crystallization was studied by means of five different sets of crystallization or MRI-monitored drying tests on plaster/substrate or painted stone specimens.

Preliminary inspections were carried out on 31 decayed buildings all over the country and allowed the selection of the following five case studies: Salvas Chapel (Sines), Alhos-Vedros tide mill (Moita), Despacho House (Pereira), Cloister of Sta. Clara-a-Nova Monastery (Coimbra) and São Sebastião church (Almada).

The five case studies were investigated by means of site inspections and a systematic sampling/testing procedure (Gonçalves *et al.* 2005, Gonçalves *et al.* 2006b). This procedure included HMC/MC profiles obtained by means of samples collected by powder drilling, as well as XRD or IC analyses on pure salt or salt-loaded samples.

Test panels of two traditional rendering systems and three industrial systems specific for salt loaded walls were made in São Sebastião church. The performance of the systems was monitored for three years (Gonçalves and Rodrigues 2005a).

The main conclusions achieved by this research are presented in the next section

7.2 Conclusions

7.2.1 - Test methods

<u>Salt crystallization tests cannot serve as general performance tests for building</u> <u>materials.</u> The key-point is that the two main aims of these tests, simulating reality and speeding up damage, are contradictory. Indeed, when damage is accelerated, distinct decay features can arise. Further, the influence of the experimental conditions may vary for different materials or boundary conditions.

Nonetheless, salt crystallization tests can be used to compare the influence of different factors on salt decay processes or to investigate salt damage mechanisms. These tests can be performed under controlled conditions and, hence, allow sound comparisons between different materials. Further, they can be designed in such a way as to isolate possible influencing factors.

Adequate crystallization tests for plasters and renders should in general use specimens composed by the plaster or render and their substrate. This is necessary because plaster/render properties depend on those of the substrate material onto which they are applied, because moisture transport and salt crystallization processes depend on the interaction between the two materials, as well as because plasters and renders do not work independently from the substrate and have a chief role of protecting the masonry.

The COMPASS test presented in Chapter 5 showed to be a good alternative to common crystallization tests. Its main limitation is that, although much faster than previous test procedures on plaster/substrate specimens, it may take a long time (in the order of six months) before five complete wet/dry cycles are achieved. However, since the salt-accumulation behaviour is probably the most fundamental feature of plaster and renders, it is always possible to reduce the number of cycles and carry out

an evaluation based on salt-distribution profiles, even when no damage has occurred yet. In any case, it is recommendable to measure the salt-distribution in, at least, one specimen of each type at the first observation of damage. These distribution profiles should be obtained, whenever possible, by means of slicing techniques involving a minimum loss of material.

<u>MRI provides a clear understanding of moisture migration in porous building materials,</u> <u>and, hence, of fundamental salt decay features.</u> It is the author's conviction that a more extensive use of MRI in this thesis would have allowed a better understanding of the salt decay processes that took place, namely, in Chapter 5 crystallization tests.

The HMC method can allow sufficiently accurate estimation of the salt content.

- Absolute salt content values can only be obtained for salts of well-known thermodynamic properties (using tables of thermodynamic data), salts available/replicable in the laboratory (using experimental correlation lines obtained with control samples) or salt mixtures of these types of salts (by either of the methods, depending on the type of salt). Particularly in the case of salt mixtures, the HMC measurements should be carried out at a high RH, so that dilute solutions are obtained where there is little interaction between the dissolved ions.
- Relative evaluation of the salt content is possible for salts of unknown thermodynamic properties, which are neither available nor are replicable in the laboratory, because the HMC is directly proportional to the salt content. This is typically the case of building samples which are normally contaminated with complex salt mixtures. However, the less homogeneous the salt mixture from sample to sample, the less reliable the HMC method.
- The accuracy of the HMC method depends on the capability of the climatic chamber to generate accurate and homogeneous environmental conditions in space and in time. Minor RH deviations may induce significant HMC variations because the HMC of soluble salts increases exponentially with the RH. Hence, the higher the actual RH of the testing environment, the higher the HMC variations produced even by small RH deviations.
- This uncertainty can be reduced by the use of control samples. Pure salt samples of the contaminant salt allow obtaining a correlation line in agreement with the actual environmental conditions in the chamber (their HMC is that of a contaminated sample with 100% salt content). Pure salt samples of a well known salt such as NaCl, whose RH_{eq} is not much affected by temperature changes, can be used to evaluate the agreement of the actual RH to the set-point RH. This is done by comparing the actual HMC of these control samples to the expected HMC. Several control samples may be spread throughout the chamber in order to evaluate the spatial homogeneity of the environmental conditions in the chamber.
- The effects of (small) RH oscillations can, if necessary, be minimized by using deep-narrow recipients. Hygroscopic equilibrium takes longer to be reached with these but the HMC fluctuation is smoothed.

 It is recommendable to perform preliminary HMC experiments, particularly in the case of poorly known salts, to ensure that the measurements are effectively carried out at an RH above the RH_{eq}^{sat} of the salt.

<u>The use of aqueous solutions for generating constant RH inside closed containers</u> (ASTM 2002, for instance) is not a safe method when salt-loaded materials are involved. Indeed, both RH and temperature may be disturbed during significant periods by secondary solutions that exist on wet materials or, when the RH is above the RH_{eq}^{sat} of the contaminant salt, form on dry materials. Therefore, these methods should in general be avoided with salt-loaded materials. They are safe only in the case of dry materials stored at RH bellow the RH_{eq}^{sat} of the contaminant salt.

7.2.2 - Salt decay processes

<u>Soluble salts may significantly influence drying of porous building materials</u>. The lower RH_{eq} of salt solutions, in comparison to pure water, induces a lower vapour pressure gradient between the material and the environment. It is also probable that the effective surface of evaporation is reduced, possibly, by deposited salt crystals. For all these reasons, when salts are present, the evaporation rate is lower and, thus, the drying front tends to be located closer to the surface. In drying tests performed on fully saturated specimens, there are longer periods of surface evaporation and higher amounts of moisture eliminated by surface evaporation.

<u>Salt-induced dampness arises from the lower drying rate of salt-loaded materials.</u> Indeed, dampness problems can occur even when the air RH is lower than the RH_{eq} of the contaminant solution because, in comparison to pure water: (i) the drying front is located on the outer surface for lower liquid supply fluxes; (ii) there is an increase in the height of capillary rise.

<u>Hygroscopicity is a particular case of this general feature and occurs when the air RH is higher than the RH_{eq} of the salt solution in the material.</u> In this case, the vapour pressure gradient is negative and, hence, water vapour diffuses from the environment to the material. Two major factors can account for the frequent severity of hygroscopic dampness: (i) the HMC of soluble salts can be very high, particularly when compared to that of building materials; (ii) very small oscillations in the RH can induce significant moisture content variations in salt-loaded masonry, particularly at a high RH.

Some salts tend to crystallize at the surface or higher on the walls, while others tend to be deposited within the pores or closer to the ground because they have a distinct influence on drying of porous materials. Indeed, the lower the evaporation rate, the higher in walls or closer to the surface crystallization tends to occur.

<u>Crystallization may take place in the masonry even when very absorbent plasters or</u> <u>renders are used</u>. Depending on the drying kinetics, local drying fronts may develop at the interfaces between materials with distinct porosity/porometry or, particularly for materials with a wide range of pore sizes, in the larger pores. The faster the drying rate of the masonry, the higher the risk of in-depth crystallization taking place. In slow drying conditions, all the pores and materials will have more time to drain the liquid solutions. Hence, less salt is expected to be deposited in the interior and more efflorescence to arise. This can happen, for example, when air temperature is low, air RH is high, wind velocity is low or when a paint layer of low vapour permeability exists.

The depth and thickness of the layer at which soluble salts accumulate in porous materials depends on the position but also on the width of the drying front. Indeed, in evaporative processes, soluble salts crystallize at the drying front. The lower the liquid/evaporation flux ratio, the deeper in the wall the drying front is located. Hence, the overall position of the front depends on the (vapour and liquid) moisture transport properties of the material, as well as on all factors that influence evaporation (such as the type of salt, solution concentration, environmental conditions, presence and type of paint layer) or the transport of liquid (for example, the moisture content in the substrate and the suction properties of the substrate material). The width of the drying front depends on the homogeneity of the pore network and on the liquid/evaporation flux ratio. The more heterogeneous the pore network (as regards both pore suction and size) and the lower that ratio, the more diffuse (thicker) the drying front.

The distribution of salt across a diffuse drying front depends on the solubility and supersaturation tendency of the salt, as well as on the concentration of the salt solution. Indeed, evaporation occurs all across a diffuse drying front and, hence: (i) the lower the solubility and the more concentrated the solution, the sooner salts will crystallize on their way across the front; (ii) for salts that may supersaturate or possess several crystal forms that precipitate at different concentrations, such as Na_2SO_4 , a more even distribution of salt can arise across the drying front.

7.2.3 - Sodium chloride and sodium sulfate

One of the main reasons for sodium chloride being typically much less damaging than sodium sulfate in laboratory crystallization tests is probably the lower vapour pressure of saturated or equally concentrated sodium chloride solutions. Due to the lower evaporation rate, a more superficial and sharper drying front occurs for sodium chloride. Since fully saturated specimens are the rule in these tests, drying Stage I will certainly be much longer for sodium chloride. Hence, a higher percentage of sodium chloride than of sodium sulfate will crystallize as harmless efflorescence. This feature is stressed when very concentrated salt solutions are used, as often happens in laboratory tests, because the vapour pressure difference between the two salt solutions is higher.

When the conditions are such that the drying front is diffuse and tends to stay inside the material, sodium chloride can cause even more severe damage than sodium sulfate. A more localized crystallization of sodium chloride is likely to be a relevant factor. Indeed, sodium sulfate tends to be distributed in a larger volume probably due to its ability for supersaturating and for precipitating at different concentrations. The discrepancy between sodium chloride's in-lab ineffectiveness and field effectiveness can be, to some extent, due to unpainted specimens being normally used in laboratory tests while in buildings painted surfaces are the norm. In fact, material damage by soluble salts in buildings often consists mostly of paint peeling, a situation that is not reproduced in laboratory tests on unpainted specimens. As seen in Chapter 5, sodium chloride was harmless to an unpainted stone but caused severe damage when that stone was painted.

<u>The new value of 95.6% for the RH_{eq}^{sat} of mirabilite which was recently proposed by</u> <u>Linnow *et al.* (2006) is supported by the results of the HMC experiments presented in section 3.5. Indeed, the random behaviour of sodium sulfate at an actual RH of around precisely 95.6% was hardly explainable in the light of a RH_{eq}^{sat} of 93.6%. But they fit with and, hence, corroborate the newly proposed value.</u>

7.2.4 - Factors that influence salt damage features

Paints can cause a more superficial deposition of salt and presence of moisture at the surface for longer periods and lower liquid fluxes, as well as an increase in the affected surface area, because they can hinder evaporation and, hence, reduce the evaporation/liquid flux ratio.

<u>Mass hydrophobic additives used in low quantity (that do not render the material a fully hydrophobic one) can enhance surface damage.</u> That happens when the environmental conditions are such that the wet front is allowed to reach the outer surface of the wall. Indeed, these additives reduce the liquid flux across the plaster probably because a part of the capillaries becomes unavailable for liquid transport. The evaporation rate is also reduced because less liquid is available at the wet front per unit time. Hence, similar amounts of moisture can take longer to be eliminated when such additives are used and larger moist areas can occur.

<u>The salt-accumulation behaviour of a certain plaster/render may differ significantly for</u> <u>distinct substrates.</u> Indeed, the substrate material can strongly condition the liquid flux into the plaster, hence, the evaporation/liquid flux ratio.

<u>The saturation state of plasters/renders also conditions their salt-accumulation</u> <u>behaviour.</u> When these materials are allowed to saturate completely, evaporation takes place on the outer surface and efflorescence may occur. The material absorption/drying hysteresis will impede this front from readily receding with decrease in the liquid/vapour flux ratio.

The salt-accumulation behaviour and related features of different plasters/renders can be affected differently by soluble salts and paint layers. For instance, as seen in Chapter 5, MEP plaster behaved as an accumulating plaster with Na₂SO₄ and, when unpainted, also with NaCl. But, when painted, MEP behaved as a transporting plaster with NaCl. However, that didn't happen for Parlumière (which was always accumulating) or for LNEC (which was always transporting). It is probable that any other factors that influence the vapour or liquid fluxes can also affect differently the behaviour of distinct plasters and renders to salt crystallization.

7.2.5 - Factors that can contribute for the worsening of salt damage

Surface damage by soluble salts can become worse after restoration interventions when the evaporation rate is reduced. This can happen due to: (i) lowering of vapour permeability, for instance, by paints or cement-based plasters/renders; (ii) reduction of the liquid flux to the drying front, for example, by partially hydrophobic plasters/renders, poorly adherent coverings or plasters/renders with a reduced suction capability from the substrate.

Indeed, in walls with rising damp: (i) the moisture content decreases with height; (ii) the water rises up to the height where the liquid supply flux from the ground is balanced by the evaporation flux through the surface of the wall. Hence, when the evaporation rate is reduced: (i) at each height, the drying front moves closer to the surface; (ii) the moist wall area is enlarged (upwards and laterally) and damage in adjacent elements may occur.

<u>Surface damage on salt-accumulating plasters may occur when low vapour</u> <u>permeability paints are used</u>. Because the evaporation rate is reduced, the drying front can move outwards, hence, can reach the outer surface and thereby enable salt-transporting behaviour.

Surface damage on salt-accumulating plasters can probably take place also when liquid continuity to the surface is established. It is reasonable to suppose that this can happen when:

- The plastering/rendering mortar is fresh. Surface evaporation can possibly persist during or after hardening, particularly in plasters/renders with strong hysteretic behaviour or under low evaporation conditions (for example, during winter, in poorly ventilated rooms or when paints are applied before the plaster is dry).
- These plaster/renders are wetted, for instance, by rain or dew point condensation. Such occurrence must, then, be prevented when salt-accumulation behaviour is to be achieved. One possibility is the use of hydrophobic paintings which, in fact, are included in most salt-accumulating systems available on the market.

<u>Carbonate salts can probably account for many of the cases where damage increases</u> <u>after restorations</u>. In Portugal, carbonate salts originating from high alkali content materials seem to be a recurrent cause of salt decay. They arise, for instance, in cement-based mortars, by reaction of the alkali hydroxides in these materials with carbonic acid formed by dissolution of air CO_2 in the water present in damp walls.

7.2.6 - Other field conditions that favour salt damage

<u>Cracking of plasters and renders can aggravate salt decay</u>. Capillary cracks enhance the transport of liquid, hence, speeding up the decay rate.

Rising damp is probably one of the main sources of moisture in old buildings. Dew point condensation can be another relevant source of moisture. It is possible that

surface sources of moisture, namely dew point condensation, are particularly effective decay agents within decay processes that take place mostly at the surface, such as carbonate formation in high alkali content materials.

7.2.7 - Diagnostic methodology

Wrong or lacking diagnoses are likely to be responsible for the failure of many restorations. Indeed, salt and moisture sources can be wrongly identified if adequate inspections and analyses are not carried out, as seen in Chapter 6. As it emerged from the interviews presented in Chapter 2, preliminary inspections or analyses by independent experts are rarely carried out neither in Portugal nor in other countries. Further, the generalized lack of scientific knowledge on salt damage processes, the unclear responsibilities and poor exchange of information between the various actors (contractors, authorities, suppliers, etc.) make it likely that ineffective diagnosis methodologies are used in many cases.

Diagnoses, to be effective, must be based both on inspections carried out by experts and on adequate sampling/testing procedures, such as those presented in Chapter 6. Expertise can provide valuable information on the causes of damage, namely as regards the sources of moisture and salt. But visual evidence is often misleading, namely, as to the true sources of moisture. Further, the type of salt, which is essential information for tracing its origin, is hardly identifiable without adequate chemical analysis.

7.2.8 - Application technique of industrial plasters and renders

<u>Plasterers must gain practical experience on applying industrial plasters and renders.</u> Well written or oral instructions are important but do not suffice to ensure a successful application.

<u>All measures needed to prevent cracking of these plasters and renders should be</u> <u>taken</u>, particularly in case of salt-accumulating or salt-blocking materials that are expected to prevent surface damage, because cracking can enhance liquid transport to the surface. It is necessary, for instance, to avoid the excess of mixing water or the application of too thick layers.

7.2.9 - Selecting plasters and renders for moist salt-loaded walls

<u>Plasters with different accumulation behaviour are adequate for different situations</u>. For instance, salt accumulating, salt blocking or moisture sealing plasters can provide surfaces free of damage. However, salt-accumulating and salt-blocking plasters are likely to reduce the evaporation rate and moisture-sealing plasters divert salt solutions to adjacent areas. Further, salt-blocking plasters do not protect the masonry from salt crystallization. Salt-transporting plasters minimize the affected area, hence providing better protection to adjacent elements. However, surface damage is more likely to arise with them.

The salt-accumulation behaviour of a given plaster/render does not depend only on the material properties. For this reason and also because crystallization tests can hardly reproduce reality, there is no straightforward answer on how to choose adequate plasters and renders for salt-loaded walls. Nonetheless, some strategies can be outlined:

- First, it is necessary to have a good understanding of the situation, not only of the physical parameters involved (type of substrate, environmental conditions, moisture load, etc.) but also of the functional requirements for the new plaster or render. As discussed above, the working principle must be appropriate to the function of the building or room, its historic value and the presence of artistic elements whose preservation is perhaps of prime concern.
- All the above discussed factors that can cause or worsen salt damage must be considered, for instance:
 - When using industrial plasters, preliminary trials should be carried out to allow the plasterers to gain practical experience with the product.
 - All measures needed to avoid plaster/render cracking and ensure a good adherence to the substrate should be taken, for instance, adequate wetting of the masonry and protection against solar radiation in the early days; see, for instance, Veiga (2004) for more information on this subject.
 - In the case of traditional (transporting) plasters, when dampness and efflorescence are to be minimized, dense or partially hydrophobic plasters or renders, as well as paints that hinder evaporation should be avoided.
 - Repaintings may enhance salt damp and efflorescence rather than reducing them because the evaporation rate is progressively lowered. Hence, if a surface requires renewal, the old paint should be previously removed.
 - Surface wetting of accumulating plasters and renders should be avoided, for example by means of a hydrophobic paint, in order to prevent liquid continuity from being established up to the outer surface.
 - In the case of salt-transporting plasters and renders increase of the drying rate or decrease of the liquid flux may cause receding of the drying front into the material and, hence, material damage by subflorescence pressure. This kind of behaviour is known to be particularly harmful in the case artistic paintings are present. It may arise, for instance, when dehumidifiers are used or capillary rise is prevented. In the last case, sacrificial layers may be used until the wall has effectively dried.
 - Though the use of hydraulic binders is typically unavoidable on moist walls, where air-lime materials can not harden properly, high alkali-content materials, such as Portland-cement mortars, should not be used because they may give rise to carbonate salts.
- Preliminary crystallization or drying tests may be used to roughly evaluate or compare the influence of different factors such as the type of substrate or the presence of paint layers. It may be more important to use realistic conditions, as regards, for example, the environmental conditions and the concentration of salt solutions, than to achieve damage in a short period of time. Salt distribution profiles

such as those presented in this thesis allow evaluation of the salt-accumulation behaviour even when damage was never achieved within the testing period. Using salts with very different properties (namely, relative equilibrium humidity and solubility) may give a better perspective of the range of possible behaviours. Sodium chloride and sodium sulfate seem appropriate but other possibilities may exist. Similarly, the use of different environmental conditions may allow the assessment of extreme behaviour, for instance, during winter or summer.

- Site tests by means of test panels should be used whenever possible. They are the best means of evaluating the true behaviour of a given plaster/render under certain conditions because the boundary conditions are the real ones. Further, site tests allow the consideration of factors related to the application technique which, as seen, can be very relevant. However, it must be taken into account that:
 - It takes considerable time and effort to properly prepare the substrate and make the test panels.
 - It is unlikely that, in most cases, significant results can be obtained after only a few months. But, as for crystallization tests, the accumulation behaviour may be accessed by means of salt distribution profiles.
 - Moisture and salts are often heterogeneously distributed in masonry walls. Hence, critical areas of the walls should be carefully chosen for the panels.
- A numerical computational model could be a powerful tool for simulating the behaviour of plasters and renders under different conditions. It is probably the only possible way of simultaneously considering the numerous complex and dynamically interrelated factors involved. Three-dimensional transport at the wall scale rather than only one-dimensional transport is necessary in order to evaluate possible damage to adjacent elements, define the appropriate dimension of the area to be plastered, as well as to allow consideration of the flux changes along the height of the wall or those caused by the spreading of the moisture into a larger area.

7.3 Future perspectives

Generally speaking, salt decay pathology is still poorly understood, as discussed in Chapter 2. There is an urgent need for research on several topics, for instance, salt damage mechanisms and influencing factors. This thesis allowed identification of the gaps in the several particular subjects which, hence, require further research.

The low evaporation rate of salt-loaded materials, for example, is not fully explainable by the lower vapour pressure of salt solutions. There is an additional lowering, which is probably due to a reduction of the effective surface of evaporation, but further research is needed to understand its causes. Deposition of salt crystals with a blocking effect is a straightforward hypothesis. Yet, there are other possibilities such as an increased contact angle between salt solutions and the porous material. A smaller effective surface of evaporation could possibly arise from a lower curvature of the menisci at the drying front. It is, for instance, necessary to understand whether the presence of solid crystals can affect that contact angle. It is extremely difficult to predict the performance of plasters and renders on salt-loaded walls. The use of an adequate computational model seems to be the best means of simultaneously considering the many (complex and interrelated) influencing factors. Efforts should, therefore, be made in the future to develop such models. Three-dimensional transport at the wall scale rather than only one-dimensional transport is necessary, as argued in the previous section.

Crystallization of carbonate salts from high alkali content materials is a frequent cause of damage in old buildings. It would be useful to develop a test or to establish threshold values for the alkali-content of building materials that allow accessing (or, at least, comparing) the tendency of plastering and rendering systems to give rise to alkali carbonate salts on moist walls. Further, a better understanding of this process is necessary, namely as regards the relevance of surface moisture sources such as dew point condensation.

Evidence was found which suggests that dew point condensation is a frequent source of moisture in old Portuguese buildings. More work needs to be carried out on selected buildings to confirm this hypothesis, as well as to evaluate the relevance within salt decay processes of dew point condensation and other surface sources of moisture such as hygroscopicity.

Cracking of plasters and renders seems to aggravate salt decay processes. Such an effect was observed in two case studies in this thesis where very high moisture contents existed in the masonry. Further research is needed to understand if and how shrinkage cracking of plasters and renders is enhanced by substrates wet in permanence or during long periods.

Paint damage is sometimes due to direct pressure by salts that crystallize at the interface with the underlying material. However, in other cases, paints peel off without any (sub)efflorescence pushing them. More research is necessary to understand which the effective damage mechanism is in this case. There are two main hypotheses: (i) vapour pressure – if the adherence of the paint to the substrate is previously weakened by salt deposition at the paint/plaster interface, detachment of paints that hinder vapour transport could more easily arise due to vapour pressure; (ii) crystallization pressure – if some salt is deposited in the porous system of the paint layer, crystallization pressure could cause dilation of such thin and elastic layer, particularly if its adherence to the substrate was already weakened by salt deposition at the interface.

When accumulating plasters or renders are applied, liquid continuity may be established up to the surface by the water in the fresh mortar. Further work is necessary to understand the relevance of such an effect as a cause of surface damage on accumulating plasters and renders. Indeed, it is possible that surface evaporation persists, during or even after hardening, in plasters/renders with strong hysteretic behaviour or under low evaporation conditions.

Further research is needed to confirm whether common drying curves provide sufficiently accurate identification of the moment when the drying front totally recedes

into the specimen. That moment is usually assigned to the critical point where the slope of the moisture-content-versus-time curve starts decreasing. Yet, when a diffuse rather than a sharp drying front occurs, there is not an immediate transition from Stage I (where evaporation takes place at the surface) to Stage II (where the evaporation front is inside the material). A transitional period exists where a decreasingly significant part of the pores close to the outer surface are involved in transporting the liquid to this surface. Since there may be a decrease in the drying rate already during that transitional period, a correct identification of the beginning of Stage II may not be possible.

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Annex I – Magnetic resonance imaging

I.1 Introduction

Magnetic Resonance Imaging (MRI) is a non-destructive imaging technique for quantitative mapping of certain chemical elements in materials. In this thesis use is made of proton NMR (Nuclear Magnetic Resonance) with the aim of monitoring the moisture content in porous building materials during drying.

The immense possibilities of NMR and MRI have already justified the attribution of two Nobel Prizes, in 1952 for Physics and in 2003 for Medicine, respectively. MRI has been used in medical applications since the 1980s, merely one decade after the first images were presented. More recently, MRI techniques have also proved to be suitable to study moisture transport in porous building materials.

In these NMR techniques, the sample is magnetized by a strong static magnetic field. In absence of a magnetic field, the nuclei in the molecules are randomly oriented due to thermal energy. Hence, the net magnetization \vec{M} , i.e., the vector sum of the magnetic moments of the individual protons, is null. The sample is then placed in a magnetic field B₀, which causes the individual magnetic moments to align and to produce a net magnetization $|\vec{M}| > 0$. In such conditions, the individual magnetic moments, as well as their resultant \vec{M} , precess in phase around the direction of B₀ at the so called Larmor frequency *f* (MHz):

$$f = \frac{\gamma \cdot B_0}{2\pi} \tag{1.1}$$

In this equation, γ (MHz.T⁻¹) is the gyromagnetic ratio of the nuclei of interest and B₀ (T) the strength of the magnetic field.

The nuclei in the sample are afterwards manipulated using radio frequency (RF) pulses by which a secondary oscillating magnetic field $B_1 << B_0$ is applied perpendicularly to B_0 . These RF pulses have to be at the Larmor frequency which is the resonance frequency of the nuclei. Because of the condition in Eq 1.1, NMR methods can be made sensitive to one specific type of nuclei. In most experiments on building materials, hydrogen protons (γ = 42.6 MHz.T⁻¹) are measured and therefore water is detected.

B₁ will cause \vec{M} to rotate away from its equilibrium position. The intensity and duration of the RF pulse determines the degree of rotation of \vec{M} .

At the termination of the RF pulse, the nuclei return to equilibrium, in a process called relaxation, by emitting electromagnetic radiation and by re-transferring the energy to the surrounding molecules. This emission of energy is observed as a NMR signal. The signal after a 90° RF pulse (a pulse which makes \overline{M} to rotate 90°) is referred to as FID (Free Induction Decay)

Here, as in general, use is made of a "spin echo sequence". In this case, the 90° RF pulse is followed by a 180° pulse resulting in a so-called spin-echo signal. The amplitude of the spin-echo signal S (a.u.) is proportional to the number of nuclei excited by the RF pulse and is given by:

$$S = \rho . e^{-\frac{TE}{T_2}} . (1 - e^{-\frac{TR}{T_1}})$$
(1.2)

In this expression, ρ is the density of the hydrogen nuclei in the sample, T_1 (s) the spin-lattice relaxation time, TR (s) the repetition time, T_2 (s) the spin-spin relaxation time and TE (s) the echo-time (Westbrook 2002, for example).

 T_1 characterizes the period of time necessary for restoring thermal equilibrium. It is a measure of the rate at which energy, which was transmitted to the system by the RF pulse, can be re-transferred from the nuclei to the lattice by random collisions between molecules. T_2 expresses how fast spin-spin interactions will cause dephasing of the protons after termination of the RF pulse. *TR* and *TE* are operational parameters. *TR* is the interval between two consecutive spin-echo sequences. A *TR* long in comparison to T_1 (*TR*~4 T_1) is used so that thermal equilibrium has time to be re-established before each new pulse sequence begins. In that case as can be seen from Eq. 1.2, T_1 has no significant influence on the intensity of the signal. *TE* is the time between the 90° excitation pulse and the echo signal acquisition. In building materials, due to the presence of magnetic impurities, T_2 relaxation is short so a short *TE* is preferred.

If a sample is placed in a uniform static magnetic field, all the protons in the sample will have the same value f of the Larmor frequency (Eq. I.1). The signal re-emitted by the relaxing nuclei will also comprise a single component, at frequency f. To allow spatial discrimination, a magnetic gradient is applied. Hence the protons in the sample will have a Larmor frequency that depends on their position. RF pulses at the appropriate frequency can then be used to measure the density of the nuclei at a given position in the sample.

In this thesis, MRI was used to monitor the drying process of specimens filled with purewater or salt solutions. Two experimental set-ups were used: one allowing one-dimensional (1D) determination of hydrogen and sodium and the other allowing two-dimensional (2D) determination of hydrogen. The experiments were performed during two stays with the group Transport in Permeable Media at the department of Applied Physics of the Eindhoven University of Technology in the Netherlands.
I.2 One-dimensional technique

The 1D MRI setup (Pel 1995) was used to monitor the drying experiments presented in section 5.6. The NMR apparatus is depicted in Fig. I.1. In this, an iron-cored electromagnet is used to produce a magnetic field of 0.78 T. Anderson gradient coils generate a constant gradient of 0.3 T/m over this main field in the vertical direction. This allows a one-dimensional resolution of the order of 1 mm in this direction. Each of the obtained NMR values corresponds, therefore, to the number of ions in a transverse section of the sample with thickness of 1 mm.



Fig. I.1 - NMR set-up for 1D measurements at TU/e composed by, from left to right: power supplies and RF transmitter, electromagnet; RF receiver; data-acquisition system

Cylindrical specimens of 20 mm diameter are used. The drying experiments were performed on plaster/brick specimens filled with pure water, NaCl solution or Na₂SO₄ solution. The specimens are, after being contaminated, wrapped in Teflon tape (Fig. I.2) in order to prevent bottom or lateral evaporation. Immediately afterwards they are inserted in the sample-holder, which is located in the middle of the NMR coil (Fig. I.3), and allowed to dry under an airflow at 0%RH (temperature \approx 18°C).

A step motor is used to move the cylindrical specimen in the vertical direction, that is, over the length of the sample, at length indents of at least 1 mm. The NMR measurements are carried out at each position. The length indents are conveniently pre-established. The smaller they are the higher the resolution but also the longer the time needed to obtain one entire profile. In these experiments, indents of 1 mm were used close to the plaster/substrate interface (typically within a 1 cm band). In the rest of the sample indents of 2 mm were used.

The results of the drying experiments are given as profiles expressing the concentration, in arbitrary units (a.u.), of the H^+ or Na^+ ion over the length of the specimen (Fig. I.4). Each profile corresponds to a different moment of the drying process. The several profiles obtained within one drying experiment give the evolution over time of the moisture content across the specimen.



Fig. I.2 – Test-specimen



Fig. I.3 – NMR RF set-up, with specimen inside the central sample-holder, and air-flow tube



Fig. 1.4 – Moisture distribution profiles obtained during one drying experiment and afterwards normalized (above). Representation of the measured specimen (below). The dashed line in the graph corresponds to an initial profile obtained in every experiment before the air flow was turned on.

RF pulses of 33 MHz and 8.9 MHz corresponding to the Larmor frequency at 0.8 T of H⁺ and Na⁺, respectively, were used. Because of the echo times used in the experiments for Na⁺ (TE=800 μ s) only dissolved nuclei are measured, that is, no signal is obtained from salt crystals. The length of the samples was approximately 50 mm in all cases which resulted in acquisition times for one entire profile of 0.5 hours for pure water, when only H⁺ was measured, or 1.5 hours for the salt solutions, when both H⁺ and Na⁺ were measured. During the drying experiments, profiles were collected every 1h for pure water and every 2h for the salt solutions. However, due to the insufficient signal-to-noise ratio achieved for Na⁺ in these samples, these profiles were not used. Therefore, only the H⁺ profiles are presented and analysed in this thesis. The operational parameters (Eq. 1.2) used for H⁺ were: TR= 1.5 sec, TE =205 μ s.

The NMR signal may be affected by magnetic impurities in the material, e.g., ferromagnetic impurities. Therefore, it was necessary to verify if the proportionality constant ρ in Eq. 4.16 varied between the two materials in a single specimen. For that purpose, the water content obtained after full immersion of brick or MEP samples in pure water during an overnight (Table I.1) was divided by the mean normalized NMR signal value obtained in the initial profile of the experiments performed with pure water on fully saturated specimens (that is, the experiment on specimen MEP5 whose results are depicted in Fig. 5.18, as well as an additional short experiment performed on MEP1 specimen). As seen in Fig. 1.5, the obtained correlation values have a limited dispersion: their maximum amplitude of variation corresponds to around 10% of the average value. Hence, the NMR signal was assumed as being proportional to moisture content regardless of the material involved.



Fig. I.5 – Ratio between moisture content and NMR signal for MEP plaster and brick

I.3 Two-dimensional technique

The 2D MRI set-up (Fig. I.6) was used to monitor the drying experiments presented in Chapter 4 and in sections 5.7 and 5.8. In these experiments, only the H^+ ion, hence water, was measured. This set-up incorporates an iron-cored electromagnet operating at a field of 0.8 T.



Fig. I.6 – NMR set-up for 2D measurements at TUE composed by, from left to right: computer; data acquisition and RF transmitter; RF receiver; electro magnet; power supplies.

Table I.1 - Water content of MEP plaster and brick after total immersion in pure water for an overnight

Water content (kg/m ³)					
MEP plaster	224				
Brick	247				

The experiments were carried out on the following types of specimens:

- Chapter 4 and section 5.8: specimens made of a lime plaster on composite substrate of the same lime mortar and stone (Fig. I.7) and filled with pure water, NaCl solution or Na₂SO₄ solution;
- Section 5.7: painted and non-painted stone specimens filled with pure water.

Immediately after contamination, the bottom face in the case of plaster/substrate specimens (their lateral faces were already sealed with epoxy resin) or bottom and lateral faces in the case of stone specimens are sealed with Parafilm M® plastic film laterally fastened with Teflon tape. The specimen is afterwards inserted in the sample-holder (Fig. I.8) which then is placed in the RF-coil of the NMR set-up (Fig. I.9). Drying proceeds under an airflow at 0%RH (temperature \approx 18°C).



Fig. I.7 – Test-specimen made of plaster on mortar/stone substrate



Fig. I.8 - Sample-holder with specimen inside



Fig. I.9 - NMR RF-coil with sample-holder connected to air-flow

The NMR set-up (Fig. I.6) uses two sets of Anderson gradient coils, i.e., in x and y directions, which allow rotating the magnetic gradient *G*. In the course of an experiment, *G* is applied at various values of the rotation angle φ (Fig. I.10) by suitably changing the values of *G_x* and *G_y*:

$$\begin{cases}
G_x = G . \cos \varphi \\
G_y = G . \sin \varphi
\end{cases}$$
(1.3)

A constant total gradient $G=0.15 \text{ Tm}^{-1}$ was used in the present case, giving a 1D resolution of around 2 mm. For each image, radial profiles are measured in 32 directions over 180° (Fig. I.10). These radial profiles correspond to projections of the nuclei density in the direction of α . They are measured by suitably changing the frequency of the RF pulse. Afterwards, by using a so-called backward projection method (program made at TUE) out of these radial profiles, the 2D distribution images are calculated.



Fig. I.10 - Representation of 2D measurement on a sample. Using variable Gx and Gy gradients, projections are measured in different α directions.

The measurement of single profile with m points (Fig. I.10) would take m^*TR . Since TR is very long (1.3 sec) this would lead to unacceptable measurement times. Therefore the profiles are measured using a so-called multiple slice spin-echo sequence. In multi-slice techniques the waiting time between spin-echo measurements at a certain position (part of TR is simply a waiting time) is used for doing measurements at a different position. By using this sequence the measurement time was cut by a factor of 4.

The NMR signal had to be averaged in order in improve the signal/noise ratio (S/N). Indeed, the signal adds linearly, while the noise, being random accumulates by a factor of the square root of n (S/N~ \sqrt{n}), n the number of measurements carried out and then averaged. Therefore, the higher n the better the S/N ratio. However, the higher n the longer the time needed for obtaining one image. In most cases, it is not feasible to have very long times for the image acquisition and a compromise has to be found. In the present experiments, the image acquisition time has to be reasonably short, when compared with the total time of the drying experiment. Hence, each measurement was repeated three times, which resulted in a total image acquisition time of 30 min with suitable S/N ratio.

During the drying experiments, images were collected every 30 min. The results are, thus, given as a sequence of 2D images (Fig. I.11) where the concentration, in arbitrary units (a.u.), of the H^+ ion is expressed by a colour scale. Each image corresponds to a given moment of drying. The backwards projection program made at TUE, by which the 2D images are obtained, allows also calculating 1D profiles, similar to those in Fig. I.4, at chosen crosssections of the specimen. Cross-sections CS1 and CS2 (Fig. I.10) were measured for some of the experiments in this thesis.



Fig. I.11 - Example of a 2D NMR image obtained during one drying experiment on a plaster/mortar/stone specimen (Fig. I.6).

Since the NMR signal may be affected by impurities, it was necessary to verify also in this case if the proportionality constant ρ in equation 4.16 varied for the different materials in the composite specimens depicted in Fig. I.7. This procedure was not applied to the specimens that included one specific type of plaster (plaster H) whose water-repellent properties changed in the course of the experiments (Fig. 5.35).

For each material, the average NMR signal value obtained at the very beginning of the drying experiments with pure water (Fig. 4.9) was correlated with the capillary saturation moisture content (Table 4.1). As seen in Fig. I.12, the correlation values were found to have limited dispersion, hence, the NMR signal could be assumed as being proportional to the moisture content regardless of the material involved. Figure I.13 depicts the MRI colour scale expressed in terms of volumetric moisture content.



Fig. I.12 - Correlation moisture content / NMR signal for plaster on mortar + stone specimens

Table I.2 - Water content of plaster L and stones D and M at capillary saturation

Water content (kg/m ³)					
Plaster L	176				
Stone D	112				
Stone M	266				



Fig. I.13 - Colour scale for the 2D NMR measurements

Annex II – Making of the test specimens for set 1 tests

All test specimens used in set 1 crystallization tests (Chapter 5) were composed of a plaster applied on brick or lime mortar substrate. Three different plasters were used: industrial plasters MEP-SP® by Strasservil and Parlumière® by Lafarge Mortiers (both on brick substrate) and traditional plaster LNEC (on each substrate). The plasters on brick substrate were either non-painted or painted with silicone paint Funcosil® by Remmers.

The specimens had flat dimensions of 40 mm x 40 mm to 50 mm x 50 mm. The thickness of the substrate was 20 mm (brick) or 30 mm (lime mortar). MEP-SP and Parlumière plasters were composed of a single 20 mm thick mortar coat. LNEC plaster was composed of an adherence coat and a base coat with thickness of 5 mm and 20 mm, respectively.

All the specimens were prepared by applying the fresh mortars on pre-wetted substrates.

MEP-SP and Parlumière specimens were prepared at LRMH within the scope of the COMPASS project. The application was carried out on half-thickness (20 mm thick) bricks by technicians from each supplier company. Specimens with flat dimensions of around 45 mm x 45 mm were later cut from the plastered bricks after removal of a 5 mm width border.

LNEC specimens were made at LNEC. The two traditional mortars that composed this plaster were prepared according to the mixing procedure in EN standard 1015-2 (CEN 1998a) using a laboratory mixer according to EN standard 196-1 (CEN 1994):

- (i) The water was introduced in the mixer. The amount of water was previously determined by means of trial mixes as that needed to achieve a flow value of 175 ± 10 mm, measured according to EN standard 1015-3 (CEN 1999a).
- (ii) The (previously mixed) dry materials were added over a period of 15 seconds with the mixer working at low speed.
- (iii) Mixing was continued for another 75 seconds at the same velocity.

Unlike MEP-SP and Parlumière, LNEC specimens had to be made individually because the traditional mortars disaggregated when cut. Hence, brick and lime mortar bases were prepared beforehand:

(i) Half-thickness (20 mm thick) bricks were cut into small blocks with flat dimensions of around 45 mm x 45 mm.

(ii) Lime mortar bases were made with flat dimensions of 50 mm x 50 mm and a thickness of 30 mm in metallic moulds. The above-described mixing procedure was followed to prepare these mortar bases. The bases cured during one month in a conditioned room at 20°C and 65%RH (demoulding was carried out at the end of the first week).

Both mortars that compose LNEC plaster were laid with the help of acrylic plaques. The plaques were placed laterally during the application and removed after around 15 minutes. After application of the adherence coat, the specimens cured during one week in the same conditioned room. Then, the base coat was applied. The specimens were kept in the same conditioned room during the following three months before testing.

Silicone paint Funcosil® was applied with a brush in a single hand when the specimens were one month old.

Plaster		Thickness	Paint	Substrate					
				Dutch red brick Pure lime mortar				ar	
				Contaminant solution					
				NaCl	Na ² SO ⁴	Pure water	NaCl	Na ² SO ⁴	Pure water
MEP-SP®		20 mm	Non- painted	S10 S11 S12	S15 S16	S7 S8 S9	-	-	-
			Silicone paint	S4 S5 S6	S17 S18	S1 S2 S3	-	-	-
Parlumière®		20 mm -	Non- painted	P10 P11 P12	P15 P16	P7 P8 P9	-	-	-
			Silicone paint	P4 P5 P6	-	P1 P2 P3	-	-	-
LNEC	Adherence coat + Base coat	ut 5 mm + 20 mm	Non- painted	LS6 LS7 LS10	LS12 LS13 LS14	LS1 LS3	LS11C LS12C LS13C	LS14C LS15C LS16C	LS9C
			Silicone paint	LS19 LS20 LS21	LS16 LS17 LS18	LS4 LS11	-	-	-

Table II.1 - Tested specimens

Annex III - Damage assessment and diagnosis form

This damage assessment and diagnosis form was compiled within the European research project COMPASS (TNO 2001). It was used for guiding the inspections carried out in the five case studies presented in Chapter 6 and has the following structure:

1 - SUMMARY

- Name of the building Location
- Type of damage decay pattern
- Materials concerned
- Tests performed
- Diagnosis
- Advice

2 - INSPECTIONS

- Date of inspection + description
- Researcher / Institute in charge of the investigation
- Reference number

3 - GENERAL INFORMATION

- Name of the building
- Address
- Owner / Responsible authority of the building
- Construction phases + date (year)
- Relevant historical calamities
- Function(s) of the building over time
- Present function (use of installations)
- Pictures of the building
- Plan of the location of the building
- Building plan

4 - STATE OF CONSERVATION OF THE BUILDING

- Type of damage and condition assessment (excellent, good, reasonable, inadequate, poor, very poor):
 - Roof
 - Façades
 - Structural elements
 - Interior
 - Floor
 - Ceiling
- Restorations or maintenance interventions performed in the past (as far as relevant):
 - Type of restorations or maintenance
 - Building part
 - Date
 - Company performing the restorations
 - Reason for restorations
 - Further information

5 - DAMAGE

- Type of damage and architectural element affected
- Location of damaged area
- Extent of damaged area (%) and depth (mm)
- Evolution of the damage
- Type of damage and material(s) concerned:
 - Wall as a whole
 - Masonry elements (brick or stone)
 - Mortar
 - (Re)Pointing
 - Render/plaster
 - Other coverings

6 - ILLUSTRATIONS

- Building plan location of the sampling
- Pictures of damaged areas

7 - ENVIRONMENT

- Climatological circumstances
- Exposition (rain, wind, etc)
- Surrounding environment (urban/rural/industrial, coastal/interior)
- Additional data

8 - DIAGNOSIS

- Hypothesis(es)
- Tests performed
- Result of tests
- Diagnosis