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Entropy parameters for falling film absorber optimization

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1	ENTROPY PARAMETERS FOR FALLING FILM ABSORBER OPTIMIZATION
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7	

8 Keywords: Entropy generation; Irreversibility; Falling film; Absorber; Horizontal tube; Optimization.

9 Abstract

A local entropy generation analysis, for water vapor absorption in LiBr-H₂O solution, is performed 10 11 referring to velocity, temperature and concentration fields obtained from the numerical solution of 12 mass and energy transport equations. The hydrodynamic description is based on Nusselt boundary 13 layer and the actual amount of irreversibility introduced is determined for a falling film over a cooled horizontal tube inside the absorber. Results are explored in different operative conditions, in order 14 15 to examine the impact of the various irreversibility sources in a wide operative range. A least 16 irreversible solution mass flow-rate can always be identified. Furthermore, a simple and general 17 thermodynamic analysis, carried out regarding a refrigerating and a heat boosting applications of absorption systems, makes evidence of a dimensionless group "Q/ σ T" that separates the weight of 18 19 the irreversibilities and gives the way to an optimization criterion applied to the absorber in order to 20 improve the whole system efficiency. Both equilibrium condition and sub-cooling of the solution at the inlet are considered for typical temperature and concentration of refrigerators' absorbers and 21 22 heat transformers' absorbers. Results suggest the importance to work at reduced mass flow-rates

23	with a thin uniform film both for refrigerators and heat transformers. In practice, tension-active				
24	additives are required to realize this condition. Also, it is highlighted that, for a fixed operative				
25	Reynolds, the two parameters defined with reference to the dimensionless group "Q/ σ T" can be				
26	maximized by specific values of the tube radius, solution sub-cooling and temperature difference				
27	between the wall and the inlet solution.				
28					
29	Nomenclature				
30	<i>a</i> Thermal diffusivity [m ² s ⁻¹]				
31	С	Molar concentration [mol·m ⁻³]			
32	Ср	Isobaric molar heat [J·mol ⁻¹ K ⁻¹]			
33	D	Mass diffusivity [m ² s ⁻¹]			
34	DA	Dimensionless absorption flux at the interface			
35	DQ	Dimensionless heat flux at the wall			
36	Ε	Entropy generation rate per tube unit length [W·m ⁻¹ K ⁻¹]			
37	g Gravity [m·s ⁻²]				
38	G	Mass flux per unit surface [kg·m ⁻² s ⁻¹]			
39	h	Molar enthalpy [J·mol ⁻¹]			
40	Н	Number of nodes in radial direction			
41	i	Specific enthalpy [J·kg ⁻¹]			
42	j	Molar flux [mol·m ⁻² s ⁻¹]			
43	k	Thermal conductivity [W·m ⁻¹ K ⁻¹]			
44	L	Mechanical power [W]			
45	т	Mass Flow-rate [kg·s ⁻¹]			
46	М	Molar weight [kg·mol ⁻¹]			

47	Ν	Number of nodes in tangential direction		
48	Ρ	Pressure [kPa]		
49	Q	2 Thermal power [W]		
50	q	Heat flux per tube unit length [kW⋅m⁻¹]		
51	r	Outer tube radius [m]		
52	R	Grid ratio in normal direction		
53	Re	Reynolds Number		
54	S Volumetric entropy generation rate $[W \cdot m^{-3}K^{-1}]$			
55	<i>s</i> Molar entropy [J·mol ⁻¹ K ⁻¹]			
56	Т	Temperature [K]		
57	и	Stream-wise velocity [m·s ⁻¹]		
58	v	Radial velocity [m·s ⁻¹]		
59	V	Total velocity [m·s ⁻¹]		
60	x	<i>x</i> Local tangential position [m]		
61	у	y Local normal position [m]		
62	Ζ	Z Tube length [m]		
63	Greek symbols			
64	Г	Mass flow rate per unit length [kg·s ⁻¹ m ⁻¹]		
65	Λ	Absorbed vapor per unit length of the tube [kg- s ⁻¹ m ⁻¹]		
66	β	Stream-wise Angle [rad]		
67	δ	Film Thickness [m]		
68	ε	Dimensionless tangential position		
69	ϕ	General parameter identification		
70	γ	Chemical potential [J·mol ⁻¹]		

71	η	Dimensionless normal position		
72	η_{Ca}	Carnot efficiency		
73	μ	Viscosity [Pa·s]		
74	θ	Dimensionless temperature		
75	ρ	Density [kg·m ⁻³]		
76	σ	Irreversibility rate [W·K ⁻¹]		
77	τ	Shear stress tensor [Pa]		
78	ω	LiBr mass concentration		
79	Subsc	Subscripts		
80	0	Standard condition		
81	А	Absorber		
82	abs	Absorption		
83	С	Condenser		
84	С	Convection		
85	СН	Chiller		
86	d	Diffusion		
87	E	Evaporator		
88	е	Equilibrium		
89	f	Friction		
90	g	Global		
91	G	Generator		
92	H2O	Water		
93	ΗT	Heat transformer		
94				

95	i,j	Node indexes
96	if	Interface
97	in	Inlet
98	min	Minimum
99	RM	Refrigerating machine
100	ТМ	Thermal machine
101	out	Outlet
102	S	Sub-cooling
103	S	Solution
104	sat	Phases equilibrium
105	t	Thermal
106	v	Vapor
107	w	Wall

108 **1. Introduction**

109 Nowadays the awareness of the environmental issues of the planet steers towards clean and efficient 110 energy conversion systems. Absorption chillers, heat pumps and heat transformers represent an 111 opportunity in this direction. The use of the vapor absorption cycle for heat driven energy systems was among the first popular and widely used methods of refrigeration. Even though the development 112 113 of vapor compression cycles has limited the implementation field of vapor absorption systems, the 114 main benefits of absorption cycle are still evident: since a negligible amount of electricity is needed, 115 waste heat can be used as the main energy source and higher reliability can be ascribed to the 116 absence of moving parts. In addition, typically used refrigerant (water or ammonia) are not responsible of ozone depletion effect. 117

118 All the real processes occurring in an energy conversion system are associated to an unavoidable degradation of the original amount of energy. The second law of thermodynamics provides a 119 120 qualitative description, which is not confined to engineering, and is critical to identify the limitations 121 imposed to a physical process. Namely, according to this general design issue, thermal design and 122 basic thermodynamics are to be employed together with the purpose of identifying the optimum size or operating regime of a certain engineering system, where by "optimum" the least exergy destroying 123 124 condition, which can still assure the fundamental engineering function, is intended. Among the 125 possible scenarios, the method of entropy generation minimization can be used to evaluate the 126 irreversibility introduced, characterize the quality of energy-conversion, and develop consistent 127 criteria for the optimization and control of the system. From this point of view, the enhancement of 128 efficiency describing the achieved technological development constitutes a secondary result of 129 entropy generation minimization [1] and entropy can be used to develop useful criteria for system 130 optimization and control.

131 From the prospective of the global system, authors in [2] and [3] have presented a literature review 132 of finite-time thermodynamics optimization of absorption refrigeration systems and analyzed the 133 various possible objective functions. In general, in order to consider the temperature level of the 134 various heat fluxes involved in the energy conversion process, a second law approach is particularly significant for the characterization of the performance of heat-driven systems. Exergy-based analyses 135 136 of water-lithium bromide absorption refrigeration systems, in both their single [4] and multiple effect configurations [5], and absorption heat transformers [6], have been performed to evaluate 137 138 performance and exergy loss of the system and its components.

The fundamental heat and mass-transfer processes constituting the absorption cycle are realized inside specific heat exchangers, whose characteristics have decisive effects on the overall system efficiency, on its dimensions and on its cost. In the conventional case of falling film heat exchangers, high transfer coefficient and low pressure drop can be obtained. However, the attempt to
experimentally and theoretically describe the complex heat and mass-transfer mechanism occurring
inside these devices is still incomplete and has not leaded to conclusive approaches. As a rule, [4],
[5], [6] and [7] have highlighted that the maximum exergy destruction occurs in the absorber and the
analysis and optimization of this device is crucial for the absorption system in refrigerator and heat
transformer applications.

In terms of modelling efforts, [8], [9], [10], [11] and [12] have discussed simplified models for falling film absorption of water vapor over a horizontal tube. Similarly, they solved the problem with a finite difference method and studied the effect of different parameters on the coupled heat and mass transfer processes.

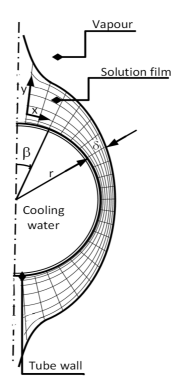
To the authors' knowledge, few studies [1], [13], [14], [15] performing a second law analysis of 152 absorption devices have been previously carried out. In particular, [16] and [17] report a second law-153 154 based analytical study for gas absorption into a laminar, gravity-driven, viscous, incompressible, 155 isothermal liquid film. The main conclusion states that entropy generation is mainly ruled by the 156 coupling effects between heat and mass transfer in proximity of the gas-liquid interface and by 157 viscous irreversibility near the solid wall. However, heat transfer and the tube wall geometry have 158 not been included in the problem. Simultaneous cooling and absorption allow the process to be 159 maintained far from the thermodynamic equilibrium at which absorption won't occur any longer.

The main purposes of this work stem from the proper expression of the local entropy generation rate representing real LiBr-H₂O absorptive films. A local analysis can be carried out by means of the numerical solution of energy and species transport equations around a cooled horizontal tube. Since entropy generation minimization has been widely accepted as a method for heat exchangers' design and optimization, a parametric study based on entropy generation analysis is performed. Finally, a simple thermodynamic analysis shows that the absorption system performance can be improved by 166 means of the optimization of the absorber design and operational regime when suitable 167 dimensionless groups, accounting for the total entropy generation introduced, are defined as 168 objective parameters.

169 **2. Model description and numerical solution**

A single horizontal tube is considered and the LiBr-H₂O solution flows viscously down over it driven by gravity as a laminar incompressible liquid, while vapor mass transfer process occurs at the film interface. The heat released by the absorption is rejected to the cooling water flowing inside the tube. The system under analysis is schematically showed in Figure 1. Heat and mass transfer characteristics of the absorber tube have been obtained by numerically solving transport of mass and energy equations, under the following main assumptions:

- 176 Steady and laminar flow without interfacial waves.
- 177 Thermodynamic equilibrium with the vapor at the film interface.
- 178 Negligible shear force between the liquid film and the vapor.
- Disturbance at the edges of the system are neglected assuming that both the tube
 circumference and length are large if compared to the film thickness.
- Physical properties are calculated with reference to the inlet values of concentration and
 temperature, but remain constant while the film is flowing on the tube.
- 183 Negligible heat transfer to the vapor phase.
- 184 Constant temperature of tube surface.
- Body fitted coordinates (x along the tube surface and y normal to it at any point) are used
- because the film thickness is small if compared to the tube circumference [12].



188

Figure 1 Local coordinate system of the flowing film

The tangential and normal velocity components (respectively, eqs. 1 and 2), from continuity and momentum equations (based on Nusselt boundary layer assumptions), are used for the numerical solution of the transport of mass and energy (respectively, eqs. 4 and 3).

192
$$u = \frac{\rho g}{\mu} \sin \beta \left(\delta y - \frac{1}{2} y^2 \right)$$
(1)

193
$$v = -\frac{\rho g y^2}{2\mu} \left[\frac{d\delta}{dx} \sin\beta + \frac{1}{r} \left(\delta - \frac{y}{3} \right) \cos\beta \right]$$
(2)

194
$$u\frac{\partial T}{\partial x} + v\frac{\partial T}{\partial y} = a\frac{\partial^2 T}{\partial y^2}$$
 (3)

195
$$u\frac{\partial\omega}{\partial x} + v\frac{\partial\omega}{\partial y} = D\frac{\partial^2\omega}{\partial y^2}$$
 (4)

196 Where the film thickness is given by eq. 5 [12].

197
$$\delta = \left(\frac{3\mu\Gamma}{\rho^2 g\sin\beta}\right)^{\frac{1}{3}}$$
(5)

- 198 The problem formulation needs to be closed by consistent initial and boundary conditions. The initial
- 199 conditions correspond to the solution temperature and concentration at the distributer, otherwise
- assuming that a complete mixing occurs, to the bulk values of the previous tube.
- 201 At x=0 and $0 < y < \delta$; T=T_{in}, $\omega = \omega_{in}$
- 202 The boundary conditions on the tube surface set a constant value of temperature and null diffusion.

203 At y=0;
$$T=T_w$$
, $\frac{\partial \omega}{\partial y}=0$

- At the interface, temperature is related to interface concentration and the heat exchanger pressure by the equilibrium condition.
- 206 At $y=\delta$; $T=T_{sat}(\omega_{if},P)$, $\omega=\omega_{if}$
- 207 The interface concentration is determined from Fick's law of diffusion:

208
$$G_{v} = -\frac{\rho D}{\omega_{if}} \frac{\partial \omega}{\partial y}$$
(6)

And the heat produced by absorption is conducted through the film towards the tube surface:

210
$$G_{v}i_{abs} = k \frac{\partial T}{\partial y}$$
 (7)

The solution approach adopted introduces a dimensionless coordinate transformation in order to map the flow domain in the physical space to a simple rectangular domain in the computational and is parallel to [18]. The dimensionless variables considered in the circumferential and radial directions are:

215
$$\varepsilon = \frac{x}{\pi r} = \frac{\beta}{\pi}$$
 (8)

216
$$\eta = \frac{y}{\delta}$$
 (9)

A cosine type grid is employed in the η direction to make the grid finer where steeper gradients are expected, i.e. near the tube wall (η =0) surface and at the interface (η =1). First order backward 219 difference is applied for the convective term in the ε direction for both energy and species transport 220 equations. In the η direction, second order central difference scheme is used both for the first and 221 the second derivative [19], respectively belonging to the convective and diffusive terms.

A second order backward difference scheme is used for the first derivative of diffusion terms of the boundary conditions at the interface. While, a second order forward difference is used for the first derivative of concentration boundary condition at the wall.

The calculation is performed between $\beta = \pi/N$ and $\beta = \pi(N-1)/N$ because of the definition domain of the velocity field (eq. 1 and 2).

3. Mathematical formulation of Entropy generation

From the calculated temperature, velocity and concentration fields, a local irreversibility analysis of water vapor absorption through a laminar, gravity driven, viscous, incompressible liquid film, flowing over a cooled horizontal tube, is performed and different entropy sources are distinguished. Owing to the simultaneous heat and mass transfer, four sources of irreversibility can be recognized and, respectively, related to heat transfer, fluid friction, coupled effects between heat and mass transfer by convection and coupled effects between heat and mass transfer by diffusion.

- 235 Physical absorption (no chemical reactions)
- 236 Steady state regime of the two dimensional flow
- 237 Newtonian and incompressible, liquid film (LiBr-H₂O solution)
- 238 Water vapor absorbed is considered as a perfect gas
- 239 The absorption process takes place at constant pressure

240 The existence of thermal, velocity and concentration gradients in the computational field 241 representing the absorptive film yields a non-equilibrium state, which is responsible of entropy variations (generally referred to as entropy generation). According to the problem formulation and
to the introduced assumptions, the volumetric rate of entropy generation due to friction, heat and
mass transfer is given by,

245
$$S_{g} = \stackrel{\mathbf{r}}{q} \cdot \stackrel{\mathbf{w}}{\nabla} \left(\frac{1}{T}\right) + \frac{\stackrel{\mathbf{w}}{\tau} : \stackrel{\mathbf{w}}{\nabla} (V)}{T} - \frac{1}{T} \stackrel{\mathbf{w}}{j_{\nu}} \cdot \stackrel{\mathbf{w}}{\nabla} (\gamma_{\nu})$$
(10)

The three terms in eq. 10 can be related, respectively, to heat transfer, fluid friction and mass transfer irreversibilities. The heat flux *q* includes the heat flux given by the Fourier Law and the enthalpy flux due to species diffusion.

249
$$q = -k_s \nabla(T) - h_v j_v$$
 (11)

As a result, a rearrangement of the expression gives eq.12.

251
$$S_{g} = \frac{1}{T^{2}} k_{S} \nabla(T)^{2} + \frac{1}{T^{2}} h_{v} j_{v} \cdot \nabla(T) + \frac{-\frac{\mathbf{u}}{\tau} \cdot \nabla(V)}{T} - \frac{1}{T} \frac{\mathbf{u}}{j_{v}} \cdot \nabla(\gamma_{v})$$
(12)

252 Considering the absorbed water vapor as an ideal gas and the only species diffusing through the liquid 253 solution, its molar concentration C_v can be derived directly from the mass concentration field of LiBr 254 (ω) resulting from the solution of energy and species transport equations.

255
$$C_{\nu,ij} = \frac{\rho_S}{M_{H_20}} (1 - \omega_{ij}) - \frac{\rho_S}{M_{H_20}} (1 - \omega_{in})$$
(13)

The molar chemical potential of the vapor can be calculated by eq. 14, as [20].

$$\gamma_{\nu}(T, P_0) = c_{p,\nu}(T - T_0) - Tc_{p,\nu} \ln\left(\frac{T}{T_0}\right) + h_{\nu,0}(T_0, P_0) - Ts_{\nu,0}(T_0, P_0)$$
257
(14)

258 Where P_0 and T_0 are the standard values of pressure and temperature, respectively, 1 atm and 298

259 K. Standard enthalpy and entropy refer to the corresponding thermodynamic state.

260 The expression of vapor molar enthalpy and entropy are, respectively, given by,

261
$$h_{v} = c_{p,v}(T - T_{0}) + h_{v,0}$$
(15)

$$s_{\nu} = c_{p,\nu} \ln\left(\frac{T}{T_0}\right) + s_{\nu,0} \tag{16}$$

263 The shear stress is given by eq. 17.

262

264
$$\tau = \mu_s \left(\frac{\partial u}{\partial y} + \frac{\partial v}{\partial x} \right)$$
(17)

265 Thus, the local volumetric entropy generation rate can be expressed as,

$$S_{g} = \frac{k_{s}}{T^{2}} \left[\left(\frac{\partial T}{\partial x} \right)^{2} + \left(\frac{\partial T}{\partial y} \right)^{2} \right] + \frac{\mu_{s}}{T} \left\{ 2 \left[\left(\frac{\partial u}{\partial x} \right)^{2} + \left(\frac{\partial v}{\partial y} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial y} \right) + \left(\frac{\partial v}{\partial x} \right) \right]^{2} \right\} + \frac{h_{v}}{T^{2}} \left[j_{v,x} \left(\frac{\partial T}{\partial x} \right) + j_{v,y} \left(\frac{\partial T}{\partial y} \right) \right] - \frac{1}{T} \left[j_{v,x} \left(\frac{\partial \gamma_{v}}{\partial x} \right) + j_{v,y} \left(\frac{\partial \gamma_{v}}{\partial y} \right) \right]$$
(18)

267 The transverse and axial molar fluxes are, respectively, given by,

268
$$j_{v,x} = C_v u - D_{vS} \frac{\partial C_v}{\partial x}$$
 (19)

269
$$j_{v,y} = C_v v - D_{vS} \frac{\partial C_v}{\partial y}$$
 (20)

By substituting the fluxes and enthalpy expressions and by calculating the derivatives of the molar chemical potential, the expression can be finally reorganized for representing the case of gas absorption into a falling liquid film.

 $S_{p} = \frac{k_{s}}{m^{2}} \left[\left(\frac{\partial T}{\partial r} \right)^{2} + \left(\frac{\partial T}{\partial r} \right)^{2} \right] + \frac{\mu_{s}}{m} \left\{ 2 \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right) + \left(\frac{\partial v}{\partial r} \right)^{2} \right] \right\} + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial v}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial u}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial u}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial u}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial u}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial u}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial u}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial u}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial u}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right] + \left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial u}{\partial r} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial r} \right] + \left(\frac{\partial u}{\partial r} \right)^{2} + \left(\frac{\partial u}{\partial$

$$= T^{2} \left[\left(\frac{\partial x}{\partial y} \right)^{2} \left(\frac{\partial y}{\partial y} \right)^{2} T^{2} \left[\left(\frac{\partial x}{\partial y} \right)^{2} \left(\frac{\partial y}{\partial y} \right)^{2} \left[\left(\frac{\partial y}{\partial x} \right)^{2} \left(\frac{\partial x}{\partial y} \right)^{2} \right] \right]$$

$$+ \left[c_{p,v} (T - T_{0}) + h_{v,0} + T \left(c_{p,v} \ln \left(\frac{T}{T_{0}} \right) + s_{v,0} \right) \right] \left[\frac{C_{v}}{T^{2}} \left(v \frac{\partial T}{\partial y} + u \frac{\partial T}{\partial x} \right) - \frac{D_{vS}}{T^{2}} \left(\frac{\partial C_{v}}{\partial y} \frac{\partial T}{\partial y} + \frac{\partial C_{v}}{\partial x} \frac{\partial T}{\partial x} \right) \right]$$

$$(21)$$

274 Different terms, related to different entropy variation sources, can be distinguished. The first term 275 of the right-hand side of eq. 21 stands for the irreversibility due to heat transfer S_t .

276
$$S_{t} = \frac{k_{s}}{T^{2}} \left[\left(\frac{\partial T}{\partial x} \right)^{2} + \left(\frac{\partial T}{\partial y} \right)^{2} \right]$$
(22)

277 The second term is due to fluid friction.

$$S_{f} = \frac{\mu_{s}}{T} \left\{ 2 \left[\left(\frac{\partial u}{\partial x} \right)^{2} + \left(\frac{\partial v}{\partial y} \right)^{2} \right] + \left[\left(\frac{\partial u}{\partial y} \right) + \left(\frac{\partial v}{\partial x} \right) \right]^{2} \right\}$$

$$(23)$$

The third and the fourth terms are related to the coupling effects between heat and mass transfer,by convection and diffusion, respectively.

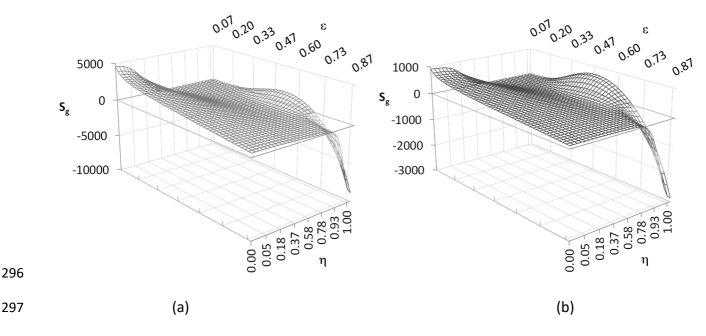
281
$$S_{c} = \left[c_{p,v}(T - T_{0}) + h_{v,0} + T\left(c_{p,v}\ln\left(\frac{T}{T_{0}}\right) + s_{v,0}\right)\right] \left[\frac{C_{v}}{T^{2}}\left(v\frac{\partial T}{\partial y} + u\frac{\partial T}{\partial x}\right)\right]$$
(24)

282
$$S_{d} = -\left[c_{p,v}(T - T_{0}) + h_{v,0} + T\left(c_{p,v}\ln\left(\frac{T}{T_{0}}\right) + s_{v,0}\right)\right]\left[\frac{D_{vs}}{T^{2}}\left(\frac{\partial C_{v}}{\partial y}\frac{\partial T}{\partial y} + \frac{\partial C_{v}}{\partial x}\frac{\partial T}{\partial x}\right)\right]$$
(25)

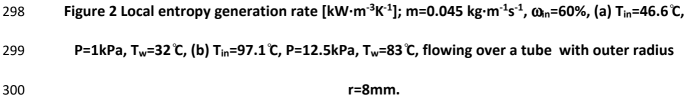
283 The solution properties are calculated for the inlet values of temperature, pressure and 284 concentration, with reference to [21].

285 **4. Results**

A general analysis has been carried out for typical conditions of a single absorber tube in a chiller and 286 in a heat transformer, respectively working with coolant temperatures of 32 °C and 83 °C, and outside 287 pressure of 1.0 kPa and 12.5 kPa. The initial conditions of the LiBr-H₂O solution film have been set at 288 289 the equilibrium temperature for a 60% concentration solution at the absorber vapour pressure. 290 Furthermore, the variation of mass flow rate due to absorption of water vapor is considered negligible. This assumption is valid for a mass flow-rate higher than 0.001kg·m⁻¹s⁻¹, and accordingly, 291 292 this analysis has been performed in a consistent mass flow-rate range [20]. Since Nusselt integral 293 solution for velocity distribution is not defined at the inlet and outlet positions, respectively $\mathcal{E}=0$ and 294 $\mathcal{E}=1$, the total entropy generation S_G has been evaluated between $\mathcal{E}=1/N$ and $\mathcal{E}=N-1/N$ inside the



295 whole film thickness.



The total entropy generation rate per unit volume S_g for an absorptive LiBr-H₂O solution flowing on 301 a cooled horizontal tube is illustrated in Figure 2 as the superimposition of the different groups 302 303 previously identified. S_g shows a local minimum in the radial direction except in the first and the last parts of the tube surface, where the total entropy generation rate is relentlessly decreasing from the 304 wall to the film interface. The minimum is determined by the contemporaneity of wall heat transfer, 305 306 friction and coupled heat and mass transfer at the interface, and it is positioned at the penetration distance of the diffusing vapor inside the film. In the second half of the tube surface the total entropy 307 308 generation rate S_g at the film interface assumes negative values, due to the only negative contribution of the group related to the coupled effects of mass convection and heat transfer S_c: this 309 310 behaviour can be explained considering that the local temperature decreases in the stream-wise direction and vapor concentration C_v increases in that region (eq. 24, where the term $C_v u/T^2 \cdot \partial T/\partial x$ is dominant).

In order to compare the performance of the system with the actual amount of irreversibility
introduced, the local volumetric entropy generation rate is integrated over the film physical domain.
Considering film thickness and the circumference of the tube the entropy generation per unit length
of the tube *E* is defined by eq. 26.

317
$$E = 2 \int_{0}^{\pi r} \int_{0}^{\delta} S dy dx$$
 (26)

The parametric analysis performed ascertains that a minimum entropy generation can always be identified in terms of the film Reynolds number (defined by eq. 27), which establishes the optimal thermodynamic condition in terms of entropy generation rate per unit length (Figure 3). As previously stated, in this work by "optimum" the least irreversible operating condition for a specified objective is meant, or otherwise, the most desirable trade-off between two or more competing irreversibilities [1]. Entropy generation minimization had been also applied to design counter-flow heat exchangers [22], [23], [24] or desiccant systems [25], [26], [27], [28].

$$Re = \frac{4\Gamma}{\mu}$$
(27)

In general, higher temperature applications have lower entropy generation and cope with higher solution Reynolds. The general trend of the different groups can be examined in Figure 3. Increasing Reynolds numbers determine increasing friction and decreasing absorption rates, and their respective entropy generation groups (E_f and E_d) show consistent trends. The combined effects of increasing extension of the entrance region, increasing film thickness and decreasing absorption heat release, globally establish an increasing behaviour of the thermal related irreversibilities E_t . The entropy generation group related to the coupled effects of mass convection and heat transfer E_c shows a local minimum (maximum of the absolute value), which can be explained considering that
when Reynolds number is increased convection is amplified while absorption heat release is reduced.
As already pointed out regarding the corresponding term of volumetric entropy generation, this
group assumes negative values.

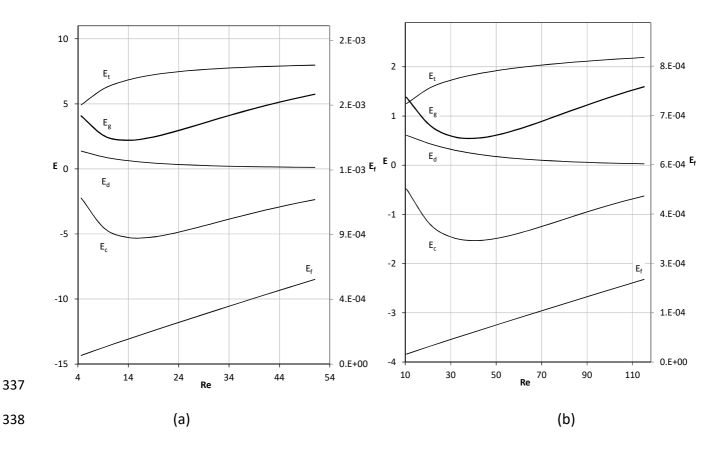
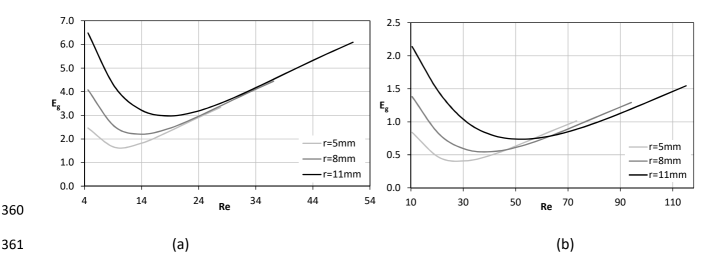
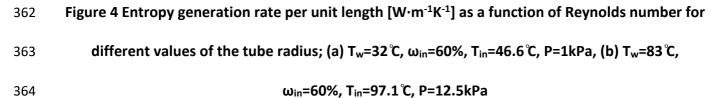


Figure 3 Various entropy generation groups per unit length of the tube $[W \cdot m^{-1}K^{-1}]$ as a function of film Reynolds number; (a) T_w=32 °C, ω_{in} =60%, T_{in}=46.6 °C, P=1kPa, (b) T_w=83 °C, ω_{in} =60%, T_{in}=97.1 °C, P=12.5kPa

Absorber design and operability parameters can be considered in the analysis. A graphical parametric study is performed evaluating the influence on the entropy generation rate of a different tube radius, coolant inlet temperature and inlet solution temperature as function of the film Reynolds number. At first, equilibrium of the inlet solution with respect to the vapour temperature (and pressure) is considered.

The influence of the radius on the global entropy generation and on each entropy generation group 347 as a function of solution mass flow rate (e.g. Reynolds Number, defined by eq. 27), for fixed inlet and 348 boundary conditions is presented, respectively, in Figure 4 and Figure 5. As a rule, lower tube radii 349 present lower entropy generation rates per unit length and match with lower solution Reynolds 350 351 values (Figure 4). The trend of the global entropy generation shows a minimum value and the 352 corresponding Reynolds number could be considered as a least irreversibility operative condition. For the chiller application case (Figure 4a), least irreversibility Reynolds numbers of 10, 14 and 23 are 353 354 obtained, respectively, for tube radii of 5, 8 and 11mm. For the same values of tube radii, but for an absorber operating at reference conditions for a heat transformer, the least irreversibility Reynolds 355 356 numbers are, respectively, 31, 42 and 52 (Figure 4b). These values have been obtained without 357 considering partial wetting of the exchange surface, but usually they correspond to solution mass flow rates which are not able to assure complete wetting, unless tension-active surfactants are added 358 359 to the solution.





Figures 5(a-b) show the effect of different tube radii on each entropy generation group. A bigger 365 radius increases the absolute value of every entropy generation group. In fact, when entropy 366 generation is integrated over the film thickness and the tube circumference the group related to 367 thermal irreversibility E_t shows a relaxingly-increasing trend with respect to Reynolds number. 368 369 Contrarily, the vapor diffusion related irreversibility E_d is characterised by a relentlessly decreasing trend. The group related to the coupled effects of mass convection and heat transfer E_c has a critical 370 influence on the global trend of the entropy generation rate per unit length Eq and, as already pointed 371 372 out, it shows a local minimum, which can be explained considering that, increasing Reynolds number, the velocity field is intensified while absorption heat release is reduced. A greater outer radius moves 373 the minimum to higher *Re* and to lower values of *E*_c. Finally, the greater the tube radius the higher 374 the friction related irreversibility E_f. 375

376 Comparing chiller and heat transformer applications, as a rule, the first corresponds to higher entropy
377 generation rate and the latter highlights a greater impact of the diffusion related irreversibility.

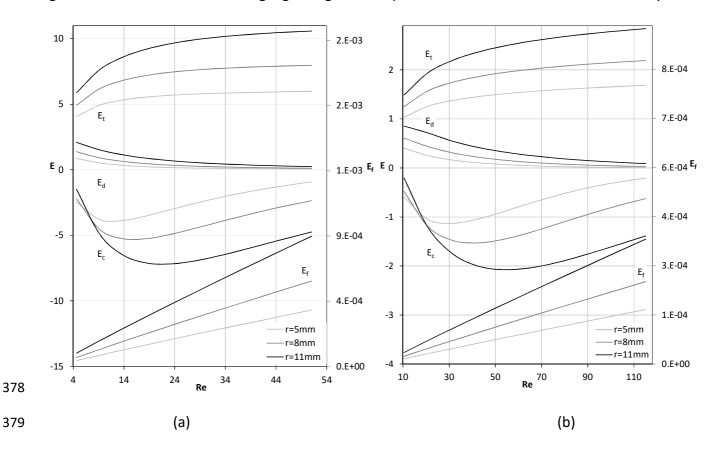
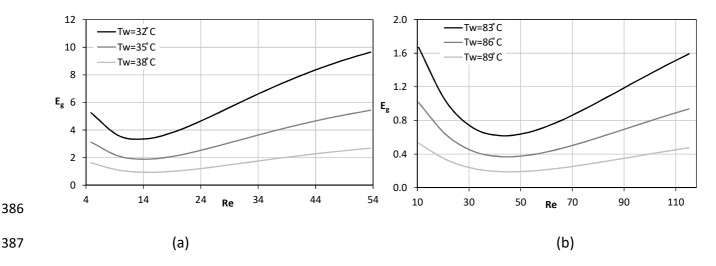
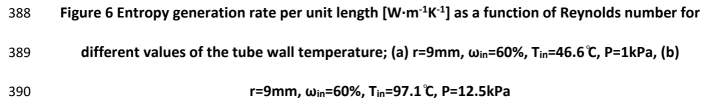


Figure 5 Effect of different radii on the various entropy generation groups $[W \cdot m^{-1}K^{-1}]$; (a) T_w=32 °C,

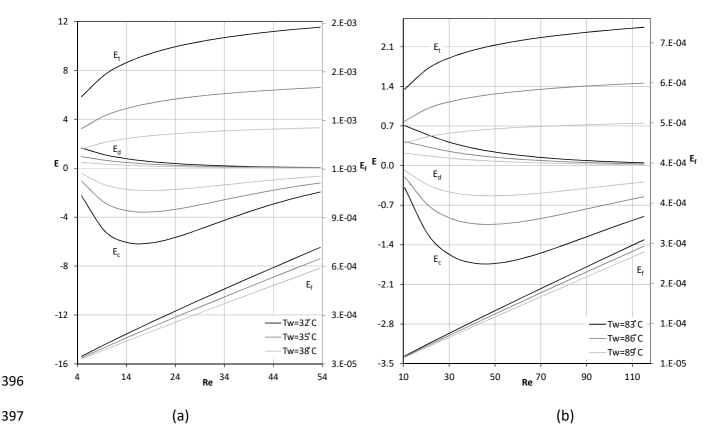
ω_{in}=60%, T_{in}=46.6°C, P=1kPa, (b) T_w=83°C, ω_{in}=60%, T_{in}=97.1°C, P=12.5kPa

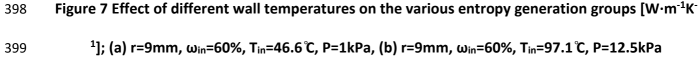
In general, a lower tube wall temperature increases heat transfer and, increasing the driving force for vapor absorption, mass transfer at the interface. Accordingly, Figure 6 makes evidence of a higher entropy generation when tube wall temperature is decreased, while the optimal Reynolds is weakly dependent on this parameter.



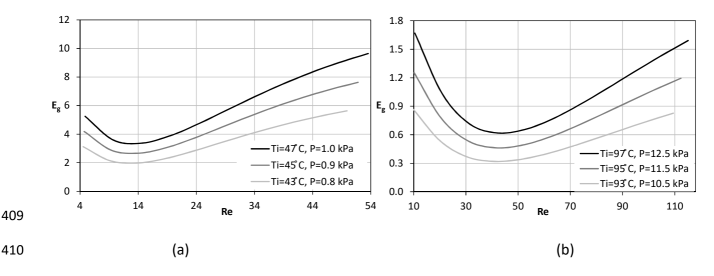


Figures 7(a-b) describe the effect of different tube wall temperatures on the various entropy generation group. A lower wall temperature increases temperature gradients and, once the temperature gradient reaches the interface, also concentration gradients. Also friction related irreversibility E_f (eq. 23) is affected indirectly by a different wall temperature which determines a different temperature field.





Furthermore, in order to analyse the effect of a different equilibrium temperature at the interface 400 (and, due to the equilibrium hypothesis, a different absorber pressure), the calculations are extended 401 402 aver a range of different inlet solution temperatures for both the chiller and the heat transformer applications. Since the tube wall temperature is maintained constant, different inlet temperatures 403 404 have a similar effect to that of different wall temperatures for a fixed inlet condition. However, for 405 the same temperature difference T_i - T_w , a change in the inlet temperature value brings about a smaller 406 entropy generation than that correspondingly obtained by changing the wall temperature. Also, 407 lower the inlet temperatures are associated to slightly lower Reynolds corresponding to the optimal 408 condition (Figure 8).





Acting again on the temperature difference T_{i} , inlet solution temperature variations have similar effects to those of a different wall temperature on the entropy generation groups as function of film Reynolds number and the corresponding graph is omitted.

416

5. Heat and mass transfer optimization

417 As a rule, the formulation of thermodynamic optimization criteria has always accompanied the 418 analysis of energy systems in order to improve their performance. Since the development of finite-419 time thermodynamics, the attention has been moved towards the concept of non-equilibrium processes and their related irreversibility sources. The use of absorption systems has attracted a 420 remarkable attention and shown a great potential in the utilization of low-grade heat sources [29] 421 [30] [31] [32]. Both chiller and heat transformer configurations have been generally analysed from a 422 first and second principle point of view [33] [34] [35] [36] [37] [38]. Based on the previous 423 thermodynamic analysis of the irreversibility associated to the absorption of vapour and the heat 424 425 transfer in a falling film configuration of the absorber, this paper aims at the definition of proper dimensionless parameters for the optimization of this component regarding the ultimate duty of the 426

system. In general, a heat transformer is a device which aims at delivering heat at a higher 427 428 temperature than the given temperature of the heat source. The component of the plant which 429 actuates this final heat transfer is the absorber. Absorption refrigeration machines are supposed to extract heat at a lower temperature than those of the ambient and the driving-fluid in the generator. 430 431 The cooling effect is realized by the evaporation of the condensed refrigerant, which is then absorbed by the solution inside the absorber, in order to be effectively pumped at the generator pressure level 432 and start the cycle again. Accordingly, the best thermodynamic condition of the absorber operability 433 434 in the two application cases considered corresponds to two different conditions.

For the first principle of thermodynamics, the COP of an absorption system can be expressed in a
general way by eq. 28 for a refrigeration application and eq. 29 in case of heat boosting one.

$$437 \quad COP_{CH} = \frac{Q_E}{L} \frac{L}{Q_G} = \frac{Q_E}{Q_C - Q_E} \frac{Q_G - Q_A}{Q_G} = \frac{Q_E}{T_C \left(\frac{Q_C}{T_C} - \frac{Q_E}{T_C}\right)} \frac{T_A \left(\frac{Q_G}{T_A} - \frac{Q_A}{T_A}\right)}{Q_G}$$
(28)

$$438 \quad COP_{HT} = \frac{Q_A}{L} \frac{L}{Q_G} = \frac{Q_A}{Q_A - Q_E} \frac{Q_G - Q_C}{Q_G} = \frac{Q_A}{T_E \left(\frac{Q_A}{T_E} - \frac{Q_E}{T_E}\right)} \frac{T_C \left(\frac{Q_G}{T_C} - \frac{Q_C}{T_C}\right)}{Q_G}$$
(29)

439 Introducing the second principle of thermodynamics, applied to irreversible cycles where isothermal
440 heat transfer without any temperature difference at the heat exchange occurs, is possible to state,

441
$$\frac{Q_C}{T_C} = \frac{Q_E}{T_E} + \sigma_{EC} \quad , \quad \frac{Q_A}{T_A} = \frac{Q_G}{T_G} + \sigma_{GA}$$
(30)

442
$$\frac{Q_A}{T_A} = \frac{Q_E}{T_E} + \sigma_{EA} \rightarrow \frac{Q_E}{T_E} = \frac{Q_A}{T_A} - \sigma_{EA} \quad , \quad \frac{Q_C}{T_C} = \frac{Q_G}{T_G} + \sigma_{GC}$$
(31)

Accordingly, the expression of the system performance in terms of COP can be rearranged highlighting the influence of components irreversibility.

445
$$COP_{CH} = \frac{Q_{E}}{T_{C} \left(\frac{Q_{E}}{T_{E}} + \sigma_{EC} - \frac{Q_{E}}{T_{C}}\right)} \frac{T_{A} \left(\frac{Q_{G}}{T_{A}} - \frac{Q_{G}}{T_{G}} - \sigma_{GA}\right)}{Q_{G}} = \frac{1 - \frac{T_{A}}{T_{G}} - \frac{\sigma_{GA}T_{A}}{Q_{G}}}{\frac{T_{C}}{T_{E}} - 1 + \frac{\sigma_{EC}T_{C}}{Q_{E}}} = \frac{\eta_{CaTM-GA} - \frac{\sigma_{GA}T_{A}}{Q_{G}}}{\frac{1}{\eta_{CaRM-EC}} + \frac{\sigma_{EC}T_{C}}{Q_{E}}}$$
(32)

446
$$COP_{HT} = \frac{Q_A}{T_E \left(\frac{Q_A}{T_E} - \frac{Q_A}{T_A} + \sigma_{EA}\right)} \frac{T_C \left(\frac{Q_G}{T_C} - \frac{Q_G}{T_G} - \sigma_{GC}\right)}{Q_G} = \frac{1 - \frac{T_C}{T_G} - \frac{\sigma_{GC}T_C}{Q_G}}{1 - \frac{T_E}{T_A} + \frac{\sigma_{EA}T_E}{Q_A}} = \frac{\eta_{CaTM-GC} - \frac{\sigma_{GC}T_C}{Q_G}}{\frac{1}{\eta_{CaHP-EA}} + \frac{\sigma_{EA}T_E}{Q_A}}$$
(33)

Broadly speaking, adopting a comprehensive point of view for chiller and heat transformer 447 application cases, eq. 32 and 33 highlight the effect of irreversibility on the system performance. They 448 449 make evidence of the importance to minimize component irreversibility, or more precisely, they 450 states that by maximizing the dimensionless groups " $Q/\sigma T$ " the efficiency of the system is improved. 451 Even though the expressions obtained for the system performance are based on a simplified approach, they make evidence of a dimensionless group, related to the irreversibilities, that indicate 452 453 a way to improve the system performance by acting also on a single device (in this case the absorber 454 of the absorption system). In a chiller or heat pump application case, the optimal performance can 455 be identified by the condition at which the absorber operates at the maximum absorption rate with 456 the least thermal power to be rejected. On the other hand, for a heat transformer, the best operative 457 condition corresponds to the maximum thermal power supplied at high temperature for the least 458 power supplied at the generator. Under this approach, based on the previous dimensionless group, 459 two different dimensionless parameters are introduced for the absorber with respect to the 460 operability of the absorption system. Eq. 34 defines the absorber dimensionless parameter used to 461 maximise heat transfer at the wall with regards to the irreversibility introduced, useful in order to 462 maximize the efficiency of the whole system in a heat transformer application.

$$DQ = \frac{q_W}{E_g T_W}$$
(34)

By maximizing the parameter DQ the thermal flux per unit length of the tube is maximized for a certain entropy generation inside the falling film. From a different point of view, at the maximum DQ, the product of the total irreversibility per unit length of the tube and the wall temperature is minimized for a certain thermal power per unit length q_W to the cooling water. The product E_gT_W embodies the thermal flux associated to the entropy variation of the whole process, in case this would be generated by heat transfer at constant temperature T_W and represents the loss of available thermal power because of the process irreversibility.

Similarly, eq. 35 defines the absorber dimensionless group used to maximise absorption at the
interface with regards to the irreversibility introduced, useful in order to maximize efficiency of the
absorption system in chiller application.

$$474 \qquad DA = \frac{\Lambda i_{abs}}{E_g T_e} \tag{35}$$

475 T_e is the equilibrium temperature of the solution at concentration ω_n (under the assumption of thermodynamic equilibrium and constant concentration at the inlet $T_e=T_{in}$) and its physical 476 significance corresponds to the temperature that would be reached if thermodynamic equilibrium 477 478 could be obtained without changes in concentration. DA is the ratio between the entropy generation 479 rate that would be produced by the heat transfer of the thermal power per unit length related to the release of the heat of absorption by the absorbed vapor at the film interface (at fixed temperature 480 481 T_e) and the global entropy generation rate per unit length E_g . Defined in this way, DA is proportional to the $Q/\sigma T$ group appearing in eq. 32. As a result, by maximizing this parameter the absorbed vapor 482 483 flux per unit length of the tube Λ (and consequently $Q_E=Z\Lambda(i_{Eout}-i_{Ein})$) is maximized for a certain entropy generation inside the falling film. Otherwise, at the maximum DA, the group $E_g T_e \cdot i_{abs}^{-1}$ is 484 minimized for a certain vapor flux per unit length of the tube. This group represents the absorbed 485 vapor flux per unit length of the tube associated to the entropy variation of the whole process, in 486

487 case this would have been generated by the release of the heat of absorption i_{abs} at constant 488 temperature T_e .

With respect to the dimensionless parameters introduced, the operability of the absorber can be investigated and then optimized, respectively, for the absorption chiller and the heat transformer application cases under analysis.

492 **5.1 Chiller**

A single value of film Reynolds number which maximizes the dimensionless group $Q/\sigma T$ (DA for the 493 chiller application, DQ for the heat transformer application) can always be identified. It can be 494 highlighted that the Reynolds at which this occurs (referred to as optimum Reynolds in the following) 495 496 is mainly dependent on the outer tube radius and on the application case (i.e chiller or heat 497 transformer operability), while a weak influence of the temperature difference between the inlet 498 solution and the wall temperature is shown (Figures 9 and 10). According to this criterion, optimal 499 Reynolds number of 9, 13 and 16 correspond, respectively, to tube radii equal to 5mm, 8mm and 11mm for the chiller application case (Figure 9b). In general, the maximum value of DA is 500 approximately constant and the film Reynolds that maximise mass transfer, with regard to DA, 501 502 increases continuously when the tube radius is increased (Figure 9a). Indeed, it can also be stated 503 that, for any fixed Reynolds there is an optimal value of the tube radius, whose value increases with 504 Reynolds, but more steeply at low than high values. Accordingly it is possible to identify an asymptotic 505 maximum value of the radius, above which is not convenient to operate with any mass flow rate.

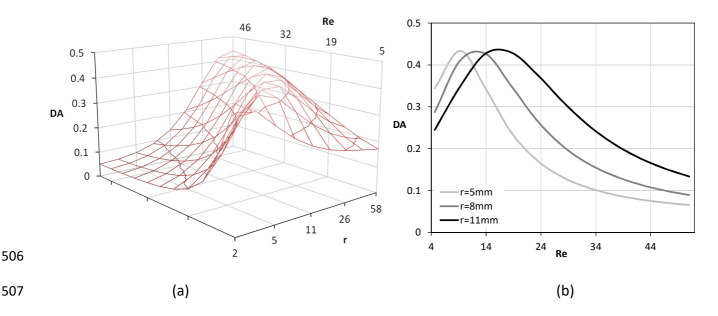
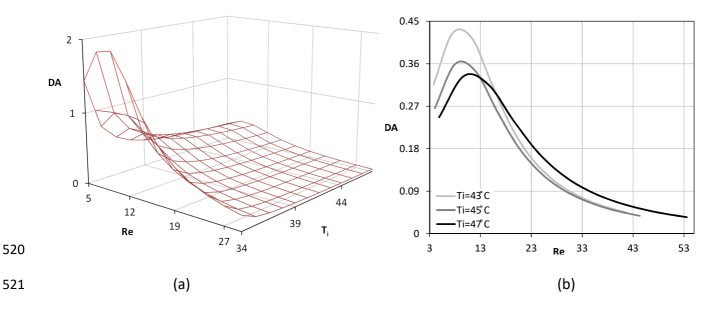


Figure 9 Dimensionless parameters DA as a function of film Reynolds number for different tube 508 radii [mm] at chiller representative conditions; ω_{in}=60%, T_{in}=46.6°C, T_w=32°C, P=1kPa (T_e=46.6°C) 509 510 On the other hand, when the difference between inlet and wall temperature is decreased (by increasing the tube wall temperature, i.e. the cooling water temperature, or decreasing the solution 511 inlet temperature), for a fixed Reynolds, the dimensionless group DA defined show a persistently 512 513 increasing trend (Figure 10a). This behaviour can be explained considering the fact that a lower temperature of the tube wall increases heat transfer (and indirectly mass transfer), but, at the same 514 515 time, the irreversibility associated to the absorption process increases at a higher rate. However, an 516 optimal condition in terms of film Reynolds number can always be identified and the corresponding 517 value increases with increasing inlet temperatures of the solution. In particular, optimal Reynolds number of 8, 9 and 11 correspond, respectively, to inlet temperatures equal to 43 °C, 45 °C and 47 °C 518 519 for the chiller application case (Figure 10b).



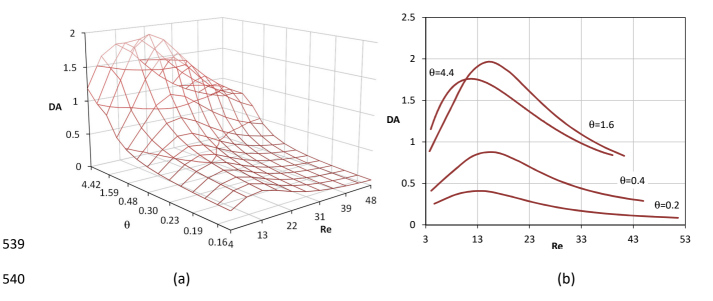
522 Figure 10 Dimensionless parameters DA as a function of film Reynolds number for various inlet 523 solution temperatures at chiller representative conditions; r=9mm, ω_{in} =60%, T_w=32.0 °C

The previous calculation has been performed under the assumption of thermodynamic equilibrium 524 525 between the inlet solution and the vapor at the absorber pressure. However, this condition, which is 526 suitable for a general irreversibility analysis, doesn't usually correspond to the operability of the absorber in actual plants [10]. Accordingly, relaxing this hypothesis, the analysis is extended to 527 528 conditions which are characterised by the sub-cooling of the inlet solution through the definition of the dimensionless parameter θ . The latter (eq. 36) expresses the temperature-boundary and initial 529 conditions in a dimensionless form, linking the sub-cooling of the solution to the temperature 530 difference between the wall and the equilibrium temperature of the inlet solution with the vapour 531 532 at the absorber pressure.

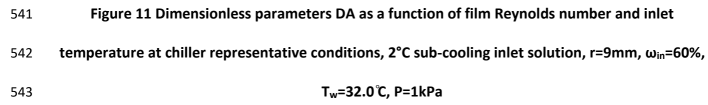
533
$$\theta = \frac{T_e - T_{in}}{T_{in} - T_W}$$
(36)

534 In general, by introducing a 2 $^{\circ}$ sub-cooling of the solution and keeping it constant, the maximum 535 values of *DA* move to higher Reynolds (Figure 11). The behaviour of the objective function *DA* with 536 respect to Reynolds and the tube outer radius in the case of a sub-cooled solution is qualitatively and

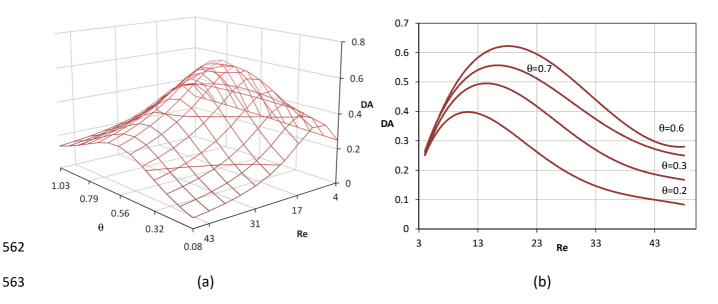
537 quantitatively similar to the case of an inlet solution at thermodynamic equilibrium with the vapor in



538 the absorber (Figure 9).



544 In order to investigate the influence of the sub-cooling, it is then possible to change the θ value 545 keeping fixed T_w and T_e value (Figure 12). Higher values of the sub-cooling at the inlet move the 546 maximum to higher Reynolds. Assuming a different standpoint in the optimization of the absorber in a chiller absorption system, we can say that a best value of the solution sub-cooling (i.e. θ) can be 547 548 identified when both T_w and T_e are fixed and furthermore, for a fixed value of θ , an optimal Reynolds 549 can always be identified, and, reversely, for a fixed Reynolds an optimal θ can be established. What is more, when the absorber works with an inlet solution characterized by the optimal value of the 550 551 sub-cooling, the corresponding optimal Reynolds is maximized (Re=17). This approach suggests again operating at low solution mass flow rates and with a small temperature difference between the inlet 552 solution and the cooling water. Consistently, in the specific case of falling film heat exchangers, the 553 554 film thickness constitutes the main heat transfer resistance with the cooling water circulating inside the tube and it has been theoretically and experimentally recognized that working with reduced solution mass flow rate can improve the system performance. As a result, operability at reduced Reynolds number is attractive for absorption plants, but the tube partial wetting at low solution flow rate needs to be considered as a critical related issue. Regarding the temperature difference between the inlet solution and the cooling water (directly related to the tube wall temperature T_w), a low value of this parameter reduces the irreversibility of the process. However, under this point of view in a specific application case the size constraint of the system is expected to be decisive.



564 Figure 12 Dimensionless parameters DA as a function of film Reynolds number and solution sub-565 cooling at chiller representative conditions, r=9mm, ω_{in} =60%, T_w=32.0°C, T_e=46.6°C, P=1kPa

Both Figures 11 and 12 show a maximum of the *DA* value. By considering the physical meaning of this parameter from a system point of view, the chiller (or the heat pump) operates at the maximum absorption rate with the least irreversibility introduced by the absorber, and, bearing in mind eq. 32, this is beneficial to the whole system COP.

570 **5.2 Heat transformer**

571 Similarly to the absorption chiller case, the absorber of an absorption heat transformer is studied by 572 means of an analysis of the dimensionless group *DQ* defined by Eq. 34. Firstly, thermodynamic equilibrium between the inlet solution and the absorbed vapor is assumed. Figures 13 shows that the maximum value of *DQ* is roughly constant, but for increasing tube radii, the optimal Reynolds number increases as well. According to this criterion, for the heat transformer application case, Reynolds number of 30, 40 and 55 maximize heat transfer in the absorber (or minimize the process irreversibility), respectively, when the tube radii are equal to 5mm, 8mm and 11mm.

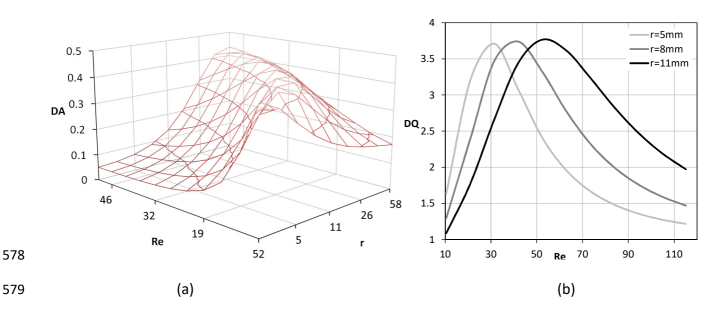
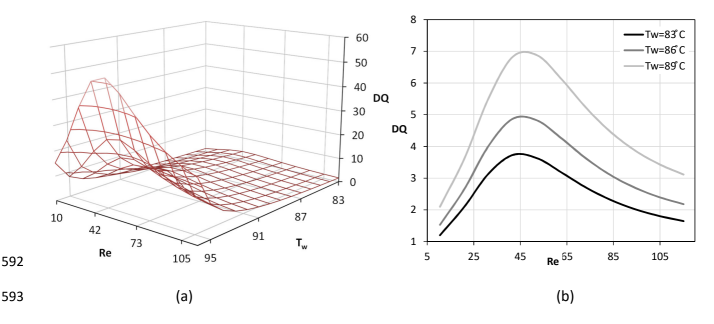


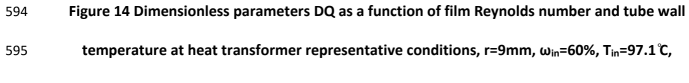
Figure 13 Dimensionless parameters DQ as a function of film Reynolds number and tube outer
 radius [mm] at heat transformer representative conditions, T_w=83.0 °C, ω_{in}=60%, T_{in}=97.1 °C,
 P=12.5kPa

Similarly to *DA*, the maximum value of *DQ* is roughly constant, but for increasing tube radii, the optimal Reynolds number increases as well (Figure 13). According to this criterion, for the heat transformer application case, Reynolds number of 30, 40 and 55 maximise heat transfer in the absorber (or minimize the process irreversibility), respectively, when the tube radii equal to 5mm, 8mm and 11mm. Moreover, the value of Reynolds which maximizes *DQ* is weakly influenced by the temperature difference between the inlet and the wall temperature, as shown in Figure 14 for different wall temperatures. *DQ* increases relentlessly when the wall temperature is augmented (i.e.

the temperature difference between the inlet solution value and the wall is lowered) for a fixed 590



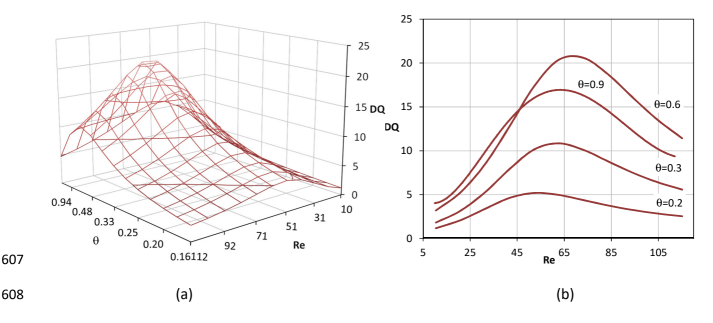
Reynolds number (Figure 14a). 591

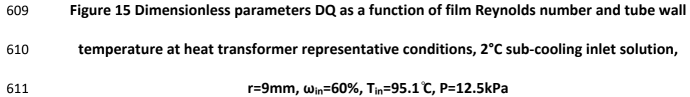


P=12.5kPa

597 When the assumption of thermodynamic equilibrium of the inlet solution is relaxed and a 2 °C subcooling is introduced consistently with the calculation performed for the chiller application, the 598 599 maxima move to slightly higher Reynolds (Figure 15). Also, the influence of the wall temperature is intensified and higher T_w are associated to increasing Reynolds. More significantly, an optimal DQ 600 condition can be shown with respect to T_w and Reynolds for the heat transformer; 0.035kg·m⁻¹s⁻¹, 601 T_w =92°C (Figure 15a). In this circumstance the optimal wall temperature is minimized and the optimal 602 603 Reynolds is maximized (Re=71). The behaviour of the previously defined objective function DQ with respect to Reynolds and the tube outer radius for a sub-cooled solution is similar to the case of an 604 605 inlet solution at thermodynamic equilibrium with the vapor in the absorber, hence, is not 606 represented herein.

596





612 When both T_w and T_e are fixed and the sub-cooling of the inlet solution becomes a parameter, a best 613 value of the latter can be identified (Figure 16). Up to that value, higher sub-cooling at the inlet move the optimal condition to lower wall temperatures and higher Reynolds. By referring to the physical 614 615 meaning of this parameter and the different operability of the absorber in the heat transformer application case considered, while maximising DQ, ceteris paribus for the operative conditions other 616 than the absorber, the whole systems operate at its highest COP and best thermodynamic state. This 617 618 condition corresponds to the maximum thermal power supplied by the heat transformer at high temperature for the least irreversibility introduced by the absorber. 619

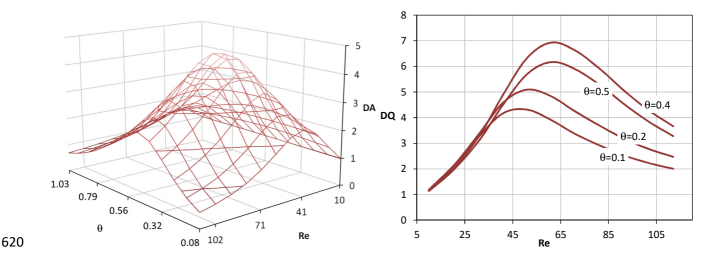


Figure 16 Dimensionless parameters DQ as a function of film Reynolds number and solution subcooling at heat transformer representative conditions, r=9mm, ω_{in} =60%, T_w=83.0 °C, T_e=97.1 °C, P=12.5kPa

624 **5. Conclusions**

The LiBr-H₂O concentration and temperature distribution inside the laminar falling film have been 625 626 obtained from the numerical solution of the coupled system of species and energy transport equations. Velocity, temperature and concentration fields, in turn, allow estimating the gradients 627 and fluxes of these variables and, finally, the local volumetric entropy generation of an absorptive 628 film flowing over a cooled horizontal tube. From the general expression obtained, various entropy 629 630 generation groups, distinguished in regard to different entropy variations sources, have been identified and investigated with respect to the critical parameters at play. This analysis characterises 631 632 the irreversibility of the process occurring in real absorbers and has been used to identify the least irreversible value of the solution mass flow rate for various operative conditions. These results make 633 evidence of the importance to work at reduced mass flow rates with a thin uniform film, and 634 635 consequently, tension-active additives are required to realize this condition.

Furthermore, a simplified and general thermodynamic analysis of the whole system performance has been proposed in order to highlight the important role of irreversibilities σ . This analysis makes evidence of a dimensionless ratio "Q/ σ T" that separates the weight of the irreversibilities. By maximizing this term the system efficiency is enhanced. Since the duty of the absorber is different in refrigeration or heat boosting applications, two different dimensionless parameters have been defined, *DQ* and *DA*, comparing, respectively, thermal flux at the tube wall and absorption at the interface with entropy generation related to the process.

643 The following main conclusion can be stated:

The parametric analysis performed makes evidence that a minimum entropy generation can
 always be identified in terms of the film Reynolds number.

The absorber of a chiller and that of a heat transformer work in different operative conditions
 which determine different irreversibility amounts introduced. In general, entropy generation
 in the heat transformer operative conditions is lower than that of the chiller and, although
 the overall trend is conserved, the minima occur at higher Reynolds.

A bigger radius increases the absolute value of every entropy generation group. Thermal 650 irreversibility E_t highlights a relaxingly-increasing trend with respect to Reynolds number. 651 Contrarily, the vapor diffusion related irreversibility E_d is characterised by a relentlessly 652 653 decreasing trend. The group related to the coupled effects of mass convection and heat transfer E_c has a critical effect on the global trend of the entropy generation rate per unit 654 length E_G and shows a local minimum, which can be explained considering that, increasing 655 656 Reynolds number, the velocity field is intensified while absorption heat release is reduced. A bigger outer radius moves the minimum to higher *Re* and to lower values of *E*_c. Finally, the 657 658 bigger the tube radius the higher the friction related irreversibility *E*_f.

659 - Comparing chiller and heat transformer applications, as a rule, the latter highlights a higher
 660 impact of the diffusion related irreversibility.

A higher difference between the solution inlet and tube wall temperatures increases both
 heat transfer and, increasing the driving force for vapor absorption, mass transfer at the
 interface. Accordingly, entropy generation is higher when this temperature difference is
 increased, while the optimal Reynolds is weakly dependent on this parameter.

When the inlet solution is at thermodynamic equilibrium with the vapor pressure in the
absorber, heat transfer at the wall and mass transfer at the film interface (represented
through the dimensionless parameters *DQ* and *DA*) can be maximised by a defined value of
film Reynolds number. It can be highlighted that the optimal value of this parameter is mainly
dependent on the outer tube radius and the application case (i.e chiller or heat transformer
operability), while a weak influence of the temperature difference between the inlet and the
wall temperature is shown.

The sub-cooling of the solution moves the occurrence of the maxima to higher Reynolds. 672 Contrarily to the case of thermodynamic equilibrium, when the temperature difference 673 674 between the inlet and the wall temperatures is changed, DQ and DA display the occurrence of an optimal condition and higher values of the sub-cooling move the maximum to lower 675 wall temperatures and higher Reynolds. Furthermore, by considering the physical meaning of 676 each parameter and the different operability of the absorber in the two application cases 677 considered, it can be observed that the optimal value of DA, at which the absorber operates 678 679 at the maximum absorption rate with the least thermal power to be rejected, corresponds to the most suitable condition for a chiller or a heat pump, while by maximising DQ in the 680 681 absorber the maximum thermal power is supplied at high temperature and, consistently, a 682 heat transformer operates at its best thermodynamic condition.

In conclusion, *DQ* and *DA* practically allow the user to realise a second law optimization of the absorber performance with respect to Reynolds, tube radius and cooling water temperature, by taking into account entropy generation. Also their influence on the overall system performance has been suggested.

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