The exergy fields in transport processes: Their calculation and use

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Abstract

This paper is a brief review of the method for analyzing the space and time dependent exergy and irreversibility fields in processes. It presents the basic equations, the method for their use, major literature sources, and three examples from the authors’ work: flow desiccation, combustion of oil droplets, and combustion of pulverized coal. Conclusions from this Second Law analysis are used to attempt process improvement suggestions. © 2005 Elsevier Ltd. All rights reserved.

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1. Exergy as an evaluation criterion

Historically, Second Law analysis in its different forms, always focusing on entropy generation and irreversibility, was at first almost entirely oriented to power generation from heat, starting with the fundamental definitions by Gouy [1] and Stodola [2] and, based on Gibbs’ work [3] expanded strongly by Keenan’s ‘availability’ concept [4,5].

Bošnjaković [6] was one of the early leaders in applying the analysis to chemical processes in his ‘fight agains irreversibility’, followed by many others including Rant, who also coined the word exergy [7], Denbigh [8], Szargut, Brodyanski, Leites, Fratzscher, Le Goff, Ishida, Rivero, Kjelstrup, and their co-workers. To illustrate some of the progress in the understanding and utility of Second Law analysis of chemical processes, the rudimentary paper by Denbigh [8] had three recommendations for keeping entropy production low, while the one by Leites et al. [9] had twelve “commandments”...
## Nomenclature

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
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<tbody>
<tr>
<td>$A$</td>
<td>exergy, kJ; Pre-exponential factor</td>
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<tr>
<td>$a$</td>
<td>specific exergy, kJ/kg</td>
</tr>
<tr>
<td>$b$</td>
<td>thickness of silica gel bed, m</td>
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<tr>
<td>$c$</td>
<td>specific heat, kJ/(kg K)</td>
</tr>
<tr>
<td>$C$</td>
<td>water vapor concentration, (kg water)/(kg mixture) ( \tilde{C} = CM_a/M_v )</td>
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<tr>
<td>$CV$</td>
<td>Control volume</td>
</tr>
<tr>
<td>$d$</td>
<td>droplet diameter, m</td>
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<tr>
<td>$D$</td>
<td>mass diffusion coefficient, m$^2$/s</td>
</tr>
<tr>
<td>$e$</td>
<td>specific exergy, kJ/kg</td>
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<tr>
<td>$E_a$</td>
<td>activation energy, kJ/mol</td>
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<tr>
<td>$E_f^{\text{av}}$</td>
<td>total (average) flow exergy, kJ</td>
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<td>$E_f$</td>
<td>local flow exergy, kJ</td>
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<tr>
<td>$f$</td>
<td>nonconservative specific body force, kN/kg</td>
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<td>$g$</td>
<td>volumetric rate of irreversible entropy production, kJ/m$^3$ K; Gibbs fuction, kJ/kg</td>
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<tr>
<td>$h$</td>
<td>specific enthalpy, kJ/kg</td>
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<tr>
<td>$H_L$</td>
<td>Sorption heat, kJ/kg</td>
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<tr>
<td>$j$</td>
<td>mass diffusion flux with respect to mass-average velocity, kg/m$^2$ s</td>
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<tr>
<td>$K$</td>
<td>kinetic energy, J</td>
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<tr>
<td>$L$</td>
<td>length of silica gel bed, m</td>
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<tr>
<td>$m''$</td>
<td>water absorption rate into silica gel, kg/(s m$^3$)</td>
</tr>
<tr>
<td>$n$</td>
<td>mass diffusion flux with respect to stationary coordinates, kg/m$^2$ s; number of species in a gas mixture</td>
</tr>
<tr>
<td>$P$</td>
<td>pressure, kPa</td>
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<tr>
<td>$q$</td>
<td>heat flux excluding radiative heat, kW/m$^2$</td>
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<tr>
<td>$q_R$</td>
<td>radiative heat flux, kW/m$^2$</td>
</tr>
<tr>
<td>$Q$</td>
<td>heat of combustion, kJ/kg</td>
</tr>
<tr>
<td>$r_i$</td>
<td>chemical species production rate, kg/s</td>
</tr>
<tr>
<td>$R, r$</td>
<td>radial coordinate</td>
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<tr>
<td>$R$</td>
<td>reaction rate</td>
</tr>
<tr>
<td>$Re_L$</td>
<td>Reynolds number = $u \infty L/\nu_f$</td>
</tr>
<tr>
<td>$Ru$</td>
<td>universal gas constant (8.314 kJ/kmol K)</td>
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<tr>
<td>$R(t)$</td>
<td>regressing droplet radius, m</td>
</tr>
<tr>
<td>$R_\infty$</td>
<td>infinity domain radius, m</td>
</tr>
<tr>
<td>$s_k$</td>
<td>specific entropy of species $k$ in the gas mixture at temperature $T$ and partial $p_k$ of component $k$ in the mixture, J/mol</td>
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<tr>
<td>$S$</td>
<td>entropy, kJ/K</td>
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<tr>
<td>$s$</td>
<td>specific entropy, kJ/kg</td>
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<td>$t$</td>
<td>time, s</td>
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<tr>
<td>$T$</td>
<td>temperature, K</td>
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<tr>
<td>$U$</td>
<td>$x$ component of velocity, m/s</td>
</tr>
<tr>
<td>$v$</td>
<td>velocity, m/s; specific volume, m$^3$</td>
</tr>
</tbody>
</table>
water content, \( \text{kg/kg} \)

flow direction coordinate, \( \text{m} \)

mole fraction of the dry air in the gas mixture

mole fraction of species \( k \) in the gas mixture

mole fraction of the water vapor in the gas mixture

coordinate perpendicular to \( x \), \( \text{m} \)

mass fraction

the axial (flow) coordinate of the RCSC

stoichiometric coefficient

unit tensor

overall heat flux excluding that by species diffusion; exergy efficiency

specific potential on a species due to any present conservative forces, \( \text{kJ/kg} \)

overall potential for all species

chemical affinity, \( \text{kJ/kg} \)

chemical potential, \( \text{kJ/kg} \)

reaction rate, \( \text{kg/(m}^3 \text{ s)} \)

porosity

mass fraction

stress tensor, \( \text{kPa} \)

the normal pressure tensor \( = P\delta + \tau_n \), \( \text{kPa} \)

density, \( \text{kg/m}^3 \)

reversible entropy flux, \( \text{kJ/Km}^2 \text{ s} \)

stress tensor, \( = \tau_n + \tau_s \), \( \text{kPa} \)

concentration of species \( i \)

pertaining to the dead state

air

chemical component of the exergy or entropy production

destruction

fluid

fuel

gas

species \( i \) index

reaction index

control volume input/output stream index

normal to surface

shear

thermo-mechanical component of the exergy

vapor

wall, i.e. the silica gel bed

free stream conditions
Bejan (cf. [10,11]) has created and motivated a large new body of knowledge linking Second Law analysis with heat and mass transfer processes, as well as with thermal power generation, refrigeration, and other fields. Some other contributions in this area, and a brief summary of the state of the art are presented below.

One of the interesting outcomes of exergy analysis is the existence of thermodynamic optima in process design, which reflect the compromises between the desirability of small temperature differences in heat transfer, and the higher flow friction and pressure drops needed to obtain them, and in chemical reactions the compromise between the increasing the driving force to accelerate reaction rates and reduce hardware costs, and the consequent undesirable entropy generation (cf. [12–14]).

It is also noteworthy that Second Law analysis is increasingly employed in numerical methods (cf. [15–18]), where the numerical formulation is improved by requiring its satisfaction at each step and in each finite element. This is often expressed as a requirement for non-negative entropy generation.

It is increasingly recognized, and included by now in practically all textbooks on thermodynamics and energy systems design (cf. [19–23]) that exergy (or Second Law) analysis must be added to the conventional energy accounting analysis during the conception, analysis, development, and design of such systems. Most of this analysis is nowadays conducted on the system level development, by evaluating the exergy values and changes of component input and output streams and energy interactions. While this can indeed identify the exergy destruction in a system component, it doesn’t deliver the detailed information about the specific process phenomena, often space and time dependent, which causes the exergy changes in it. The phenomena may include heat transfer, mass transfer, fluid mechanics, chemical and/or nuclear reactions, and the presence of fields such as gravitational, electric and magnetic. This type of detailed analysis, which we shall call ‘intrinsic’, due in the second phase of system development, is invaluable in accelerating the evolution of the innovative systems needed to meet the difficult demands of the coming century, and is the focus of this paper.

Although the objectives of exergy or entropy analysis appear to be obvious, it is very noteworthy to recall that the most important and useful ones are: (1) identification of the specific phenomena/processes that have large exergy losses or irreversibilities, (2) understanding why these losses occur (3) evaluation how they change with any changes in the process parameters and configuration, and (4) as a consequence of all these, suggestions on how the process could be improved. A surprisingly large fraction of the publications in this field deal only with (1) above, and are therefore of little use, at best.

We start by presenting the basic field equations needed for the intrinsic exergy analysis, briefly survey the state of the art, and present specific examples from our work on convective desiccation and on combustion of oil droplets and of pulverized coal.

2. The exergy/entropy production field equations

The general transient rigorous partial differential equations for computing the intrinsic irreversibilities and exergy destruction in general, needed for this level of analysis, were developed and presented by Dunbar, Lior, and Gaggioli [24], based on earlier work by Hirschfelder et al. [25], Obert [26], and Gaggioli [27,28]. Some of the key equations from the first reference are summarized below.
The reversible entropy flux is

$$\sigma = \frac{\varepsilon}{T} + \sum_i j_i S_i \quad (1)$$

where

$$\varepsilon = q + q_R - \sum_i j_i h_i \quad (2)$$

The exergy is defined as

$$a = K + \Phi + (T - T_0)S - (P - P_0)v + \sum_i (\mu_i - \mu_{i0})Q_i \quad (3)$$

where the terms on the right hand side of the equation are the kinetic, potential, thermal, strain, and chemical exergy components, respectively. The transient rate of change of exergy of a fluid element moving through the domain of interest is obtained by the substantial derivative, so Eq. (3) becomes

$$\frac{Da}{Dt} = \frac{DK}{Dt} + \frac{D\Phi}{Dt} + \frac{D[(T - T_0)S]}{Dt} - \frac{D[(P - P_0)v]}{Dt} + \frac{D[\sum_i (\mu_i - \mu_{i0})Q_i]}{Dt} \quad (4)$$

To allow detailed calculation of the individual causes for exergy change, the terms in Eq. (4) were decomposed into their basic components:

The transport of the kinetic exergy

$$\frac{DK}{Dt} = -\frac{1}{\rho} \left[ \nabla \cdot (\tau \cdot v) \right] + \frac{1}{\rho} \sum (n_i \cdot f_i) - \frac{1}{\rho} \sum (n_i \cdot \nabla \phi) - \frac{1}{\rho} \left[ v \cdot (\nabla \cdot \pi) \right] + \frac{1}{\rho} \sum (j_i \cdot f_i) + \frac{1}{\rho} \sum (j_i \cdot \nabla \phi) \quad (5)$$

has seven components, due to: (i) shear stresses, (ii) work done by nonconservative body forces on the species flux, (iii) reversible interconversion between kinetic and potential exergy, (iv) reversible interconversion between kinetic and strain exergy, (v) irreversible conversion of kinetic exergy due to friction (this part is converted to heat), (vi) irreversible conversion of nonconservative body force action on the species flux, (vii) the irreversible portion of the potential to kinetic exergy conversion.

The transport of potential exergy,

$$\frac{D\Phi}{Dt} = -\frac{1}{\rho} \sum [\nabla \cdot (\Phi j)] + \frac{1}{\rho} \sum (n_i \cdot \nabla \phi) - \frac{1}{\rho} \sum \phi f_i \quad (6)$$

has three components, due to (viii) diffusion, (ix) reversible interconversion between kinetic and potential exergy, (x) potential exergy change associated with reactions.

The transport of thermal exergy,

$$\frac{D[(T - T_0)S]}{Dt} = -\frac{1}{\rho} \left[ \nabla \cdot [(T - T_0)s] \right] + \frac{1}{\rho} S \frac{DT}{Dt} + \frac{1}{\rho} \left( \mathbf{a} \cdot \nabla T \right)$$

$$+ \frac{1}{\rho} \left( \frac{T_0}{T} \right) \left[ -\left( \tau_s \cdot \nabla v \right) - \left( \tau_n \cdot \nabla v \right) - \sum (j_i \cdot f_i) + \sum (j_i \cdot \nabla \phi) - \sum j_i \dot{R}_i - \sum (j_i \cdot \mu) \right] \frac{DT}{Dt} \quad (7)$$

has ten terms, due to (xi) heat diffusion, (xii) temperature changes, (xiii) heat conduction, the reversible interconversions between (xiv) shear strain, (xv) normal strain, (xvi) nonconservative potential
gradients, (xvii) body forces, (xviii) chemical reactions, (xix) chemical potential gradients, and (xx) changes in the reference (dead state) temperature.

The transport of strain exergy

\[ \frac{D}{Dt} \left[ -(P - P_0) \mathbf{v} \right] = -\frac{1}{\rho} \left[ \nabla \cdot (\mathbf{\pi}_s - P \mathbf{\delta}) \mathbf{v} - \mathbf{v} \frac{DP}{Dt} - \frac{1}{\rho} \left[ \nabla \cdot (\mathbf{\tau}_s) \mathbf{v} - \mathbf{\tau}_s : \nabla \mathbf{v} \right] + \mathbf{v} \frac{DP}{Dt} \right] \]

has five terms, due to (xxi) net transport of strain exergy via normal stresses, (xxii) and (xxiii) are reversible interconversions between normal strain and kinetic exergy, (xxiv) irreversible conversion of strain exergy, and (xxv) changes in the reference (dead state) pressure.

The transport of chemical exergy:

\[ \frac{D}{Dt} \left[ \sum_i (\mu_i - \mu_{i0}) \chi_i \right] = -\frac{1}{\rho} \sum_i [\mathbf{v} \cdot (\mu_i - \mu_{i0}) \mathbf{j}_i] + \sum_i \chi_i \frac{d\mu_i}{dt} + \frac{1}{\rho} \sum_i [\mathbf{v} \cdot (\mu_i - \mu_{i0})] - \frac{1}{\rho} \sum_i \chi_i \mathbf{R}_j - \sum_i \chi_i \frac{D\mu_{i0}}{dt} \]

has 5 components due to (xxvi) chemical exergy diffusion, (xxvii) reversible interconversions with thermal and/or strain exergy as can be understood from the Gibbs–Duhem Eq. (10)

\[ \sum_i \chi_i d\mu_i = vdP - sdT, \]

(xxviii) and (xxix) irreversible conversion of chemical to thermal exergy by diffusion and chemical reactions, respectively, and (xxx) changes in the reference (dead state) chemical potential.

The breakdown of exergy into such detailed components allows a most careful and detailed examination of the source of irreversibility and exergy destruction. Many of the components may in specific applications be negligible, but we can not know that before they are calculated. For example, strain exergy may be negligible compared to thermal exergy in many cases, but will dominate in cases where the velocity and velocity gradients are high while the thermal exergy is low.

While exergy is a very useful property, ultimate decisions on process improvement are most often made based on economic considerations. In fact, lower exergy efficiencies are then acceptable if the overall product cost is reduced thereby. The link between exergy analysis and economics, sometimes named exergoo-economics, is under continuous development (cf. [21]), but is based on component level analysis. Proposals were made lately to assign an economic cost to local exergy values and to thus evaluate the exergy cost fields as a continuum [29,30]. It is not difficult to add a cost multiplier to exergy fields computed by the equations above, and it even may be of some use in evaluating some specific problems, but, as opposed to thermodynamic and transport phenomena and properties, costs do not have universal and commonly defined values. Furthermore, the assumption made that all components of the local exergy have the same specific cost are too simplistic and usually incorrect. Consequently, the proposed ‘Local Exergy Cost Theory’ should at best be regarded as local to the user and the specific application, and not to spatial position.

3. The information needed for rigorous intrinsic exergy analysis

Observation of these equations shows that knowledge of the field parameters \( \mathbf{v} \) (and hence \( \mathbf{\tau}, P \) (and hence also \( \mathbf{\tau}_n \)), \( T \) (and hence \( \mathbf{\epsilon} \) and \( \mathbf{\sigma} \)), \( \Omega \) (and hence also \( \mathbf{J}_i, \mu \)), of the state equations for the materials
used, to determine the thermophysical properties \((\rho, \mu, S)\), and of the chemical reactions involved (to determine \(\lambda, R, r_i\)) allows the computation of the exergy, exergy destruction, entropy, and entropy generation fields. All of these field parameters are obtained from the solution of the full field and state equations, consisting of the Navier-Stokes, energy, species conservation, entropy generation, and thermodynamic properties equations, combined in a combustion process with the reaction kinetics equations, all tightly coupled.

Naturally, at least an order of magnitude evaluation of the different components of exergy should be performed as a first step, to determine whether they can be ignored or need to be calculated more precisely.

4. Exergy analysis in heat transfer

**Forced convection.** Bejan [31] was perhaps the first to examine entropy generation in convective heat transfer, noting that this leads to an optimization opportunity between the reduction of entropy generation by using small temperature differences, and the pumping power needed. He developed expressions for the optimal diameters of cylinders in internal and external convection, which minimize entropy production. Several other studies followed along these lines, such as for external flows: across cylinders with mixed forced and natural convection [32,33], and shear flows [34]; for internal flows including examination of duct cross section shape [35,36] and swirling flows [37], and general convection as well as convection-radiation interaction in porous media [38,39].

An appropriate application of the exergy analysis results is that by Ibanez et al. [40], who analyzed some simple cases to find that asymmetric cooling (more from one side than the other) leads to entropy minimization.

**Natural convection.** Several authors have examined entropy generation in natural convection (cf. [32,33,41,42]) and at the onset of natural convection [43].

5. Heat exchange

Intrinsic exergy (or entropy generation) analysis is very useful in determining the conditions at which heat or mass exchange between bodies or streams can be performed with minimal entropy production. It was found [44,45], that this minimum is attained in exchangers that allow constant entropy production in all of their parts (equi-partition of entropy production), a condition approximated well by imposing equi-partition of driving forces along the exchanger.

6. Exergy analysis in combined flow, heat and mass transfer

Notable are the papers [46,47].

**An example: convective desiccation.** An exergy analysis of the water vapor adsorption process in a desiccant-air stream system, for laminar humid air flow over a desiccant flat bed, as well as in a desiccant-lined channel, and for turbulent humid air flow in such a channel, for different turbulence
intensities was conducted [48], and a brief summary of the major steps of the approach and of the major results is given below.

The physical system considered (Fig. 1) is a flat silica-gel-packed desiccant bed of length $L$ with a uniform air stream passing over it in parallel.

6.1. Derivation of the flow exergy of the humid air stream

Using the system of Eqs. (1)–(4) Eq. (5)–(10) for this case produces the following equation expressing the specific flow exergy on mass basis, (in kJ/kg)

$$ e_f(x, y) = T_o \left\{ \frac{c_{pa} + \tilde{C}(x, y)c_{pv}}{1 + \tilde{C}(x, y)} \left[ \left( \frac{T(x, y)}{T_o} \right) - 1 - \ln \left( \frac{T(x, y)}{T_o} \right) \right] + R \ln \left( \frac{p(x, y)}{p_o} \right) \right\} $$

$$ + RT_o \frac{1}{1 + \tilde{C}(x, y)} \left[ (1 + \tilde{C}(x, y)) \ln \left( \frac{1 + \tilde{C}^o}{1 + \tilde{C}(x, y)} \right) + \tilde{C}(x, y) \ln \left( \frac{\tilde{C}(x, y)}{\tilde{C}^o} \right) \right] $$

(11)

Then, the total volume-average flow exergy (in kJ) is

$$ E_f = \frac{m_f}{L(h - b)} \int_0^{L} \int_0^{(h - b)} e_f(x, y) dx dy $$

(12)

where $m_f$ is the mass of the moist air in the flow region and it is calculated by

$$ m_f = \rho_f (h - b) L $$

(13)

where $\rho_f$ is the fluid (air-water vapor) density, $h$ is the practical height, $b$ is the desiccant bed thickness, and $L$ the bed length.

The local flow exergy Eq. (11) in kJ in the flow region is

$$ E_f(x, y) = e_f(x, y) m_f(x, y) $$

(14)

---

Fig. 1. The physical model configuration for the flat desiccant bed.
From Eqs. (11)–(13), the volume-average thermo-mechanical and chemical components of the exergy, $E_{f, \text{th}}$ and $E_{f, \text{ch}}$ respectively, are

$$E_{f, \text{th}} = \frac{m_f}{L(h-b)} \int_0^{L} \int_0^{L} T_o \left\{ c_{pv} + \tilde{C}(x,y) \frac{c_{pv}}{1 + \tilde{C}(x,y)} \left[ \left( \frac{T(x,y)}{T_o} \right) - 1 - \ln \left( \frac{T(x,y)}{T_o} \right) \right] + R \ln \left( \frac{p(x,y)}{p_o} \right) \right\} \, dx \, dy$$

(15)

$$E_{f, \text{ch}} = \frac{m_f}{L(h-b)} \int_0^{L} \int_0^{L} R T_o \frac{1}{1 + \tilde{C}} \left( x, y \right) \left[ (1 + \tilde{C}(x,y)) \ln \left( \frac{1 + \tilde{C}}{1 + \tilde{C}(x,y)} \right) + \tilde{C}(x,y) \ln \left( \frac{\tilde{C}(x,y)}{C^o} \right) \right] \, dx \, dy$$

(16)

### 6.2. Derivation of the flow exergy of the water vapor in the desiccant bed

The amount of air that is adsorbed by desiccant is small compared to the amount of water vapor, therefore the corresponding term in Eq. (9) can be assumed to be zero. Also, a term is added to represent the heat of adsorption added to the system due to absorption rate $m^m$. Applying these conditions, the specific flow exergy is

$$e_{f,s}(x,y) = T_o \left\{ c_{pv} \frac{\tilde{C}(x,y)}{(1 + \tilde{C}(x,y))} \times \left[ \left( \frac{T(x,y)}{T_o} \right) - 1 - \ln \left( \frac{T(x,y)}{T_o} \right) \right] + R \ln \left( \frac{p(x,y)}{p_o} \right) \right\}$$

$$+ RT_o \frac{\tilde{C}(x,y)}{(1 + \tilde{C}(x,y))} \left[ \ln \left( \frac{1 + C^o}{1 + \tilde{C}(x,y)} \right) + \ln \left( \frac{\tilde{C}(x,y)}{C^o} \right) \right]$$

$$- \left( \frac{H_l m^m(x,y)}{\rho_f} \right) \cdot \left( 1 - \frac{T_o}{T(x,y)} \right) \Delta t$$

(17)

where $H_l$ is the adsorption heat, $m^m$ is the water adsorption rate, $\rho_f$ is the fluid density, and $\Delta t$ is the time step. Then, the total volume-average humid air exergy (in kJ) in the solid desiccant is

$$E_{f,s} = \frac{m_{f,s}}{Lb} \int_0^b \int_0^L e_{f,s}(x,y) \, dx \, dy$$

(18)

where $m_{f,s}$ is the mass of the fluid in the solid desiccant bed calculated by

$$m_{f,s} = \sigma \rho_f b L$$

(19)

where $\sigma$ is the bed porosity.

The local flow exergy is also computed in kJ at any location in the solid desiccant bed from Eq. (17) by

$$E_{f,s}(x,y) = e_{f,s}(x,y)m_{f,s}(x,y)$$

(20)
From Eqs. (18)–(20), the volume-average thermo-mechanical, chemical, and adsorption heat in the desiccant components of the exergy, respectively, are

\[
E_{f, \text{th}} = \frac{m_{f,s}}{Lb} \int_0^b \int_0^L T_o \left\{ c_{pv} \frac{\tilde{C}(x,y)}{1 + \tilde{C}(x,y)} \left[ \left( \frac{T(x,y)}{T_o} \right) - 1 - \ln \left( \frac{T(x,y)}{T_o} \right) \right] + R \ln \left( \frac{p(x,y)}{p_o} \right) \right\} \text{d}x \text{d}y
\]

(21)

\[
E_{f, \text{ch}} = \frac{m_{f,s}}{Lb} \int_0^b \int_0^L \left\{ R T_o \frac{\tilde{C}(x,y)}{1 + \tilde{C}(x,y)} \left[ \ln \left( \frac{1 + \tilde{C}^o}{1 + \tilde{C}(x,y)} \right) + \ln \left( \frac{\tilde{C}(x,y)}{\tilde{C}^o} \right) \right] \right\} \text{d}x \text{d}y
\]

(22)

\[
E_{f, \text{ad}} = \frac{m_{f,s}}{Lb} \int_0^b \int_0^L \left\{ \frac{H_{i,m}^{m}(x,y)}{\rho_f} \left( 1 - \frac{T_o}{T(x,y)} \right) \Delta t \right\} \text{d}x \text{d}y
\]

(23)

It is noteworthy that the solid desiccant (without its vapor and air content) also has exergy, but it was not considered in this study because it depends on the temperature primarily and its variation over the small temperature range considered here is very small.

6.3. Computation and some results

The exergy was calculated from Eqs. (11), (15)–(17), and (21)–(23), using the velocity, temperature and concentration field results in the humid air flow and the desiccant, described briefly above and in more detail in [49].

The total exergy in the desiccant bed is about 25-fold lower than that in the humid air stream, and the overall average exergy in the combined desiccant-air stream system is dominated by the average chemical exergy in the air stream, so the values are nearly equal to those of the air stream. Typical results are shown in Fig. 2: most of the water absorption takes place in the first cm or so, where the corresponding release of heat of adsorption composes most of the exergy, and the chemical exergy component dominates downstream, where the adsorption diminished significantly.

One of the exergy analysis results shown above indicates that the release of heat during the adsorption process expends exergy without any benefit. As a matter of fact, the resulting heating of the air stream being dehumidified is most often detrimental to the ultimate use of this air, such as in airconditioning applications. Desiccant dehumidification analyzed here is generally an exergy efficient process, only < 7% is destroyed due to unused heating.

Fig. 3 shows that the exergy for the laminar flow at \( t = 4 \) s is about 9% larger than that for turbulent flow, due to the reduction in the water vapor concentration in the laminar flow model since the dominant component in the desiccant-air system is the chemical component in the air stream (Eq. (16)). Therefore, the overall average exergy decreases as the water vapor concentration increases with the time. The exergy reduction for laminar flow is about 19.3%, and it is 26.5% for turbulent flow, as the time goes from 0 to 1 s. While turbulence increases exergy reduction due to its dissipative nature, the main reason for the observed exergy reduction here is the increased mass transport rates due to the turbulent flow.
Fig. 2. (a) Exergy components, (b) water concentration, and (c) water adsorption rate distributions along the bed in the humid air stream and the desiccant bed. $u_\infty = 0.1 \text{ m/s}, Re_L = 333, T_\infty = 30 ^\circ \text{C}, C_\infty = 0.0276 \text{ kg/kg}, C_p = 0.0075 \text{ kg/kg}, T_o = 25 ^\circ \text{C}, C_o = 0.00992 \text{ kg/kg}, m_{f,s} = 1.9 \times 10^{-5} \text{ kg}, m_f = 0.0028 \text{ kg}, b = 0.00321 \text{ m}, L = 0.05 \text{ m}, h = 0.05 \text{ m}, \sigma = 0.1.$
It was also found that increasing the turbulence intensity 10-fold, from 1 to 10%, results in a decrease of the exergy by about 5% at $t = 4$ s. As the time goes from 0 to 1 s, the exergy reduction for 1% turbulent intensity is about 26.5%, and it is 30.3% for 10% turbulent intensity.

Conclusions and possible constructive uses of the exergy analysis of flow desiccation

- In laminar flow, a total of $\sim 20\%$ of the humid air exergy is expended in its drying, but 90% of this is a change in chemical exergy, needed anyway; the temperature rise is $< 2$ °C.
- In the desiccant, practically all of the exergy reduction is due to the release of absorption heat, and raising the solid temperature; desiccants with small heat of adsorption are thus desirable.
- For the same inlet velocity, a desiccant-lined channel is more effective for dehumidification than a flat bed, and proportionally $\sim 20\%$ more exergy is expended.
- Turbulent flow improves dehumidification and proportionally increases exergy expenditure 27% for $TI = 1\%$ and to 30% for $TI = 10\%$.
- Desiccant dehumidification analyzed here is generally an exergy efficient process, only $< 7\%$ is destroyed due to unused heating.
- Residence times beyond 1.5 s, and beds longer than 1 cm are detrimental.
- Since most of the absorption/desorption takes place near the desiccant entrance and rapidly decreases, optimal design may result from beds which are thicker at the entrance and are gradually thinned with $x$.
- Synergistically with improved mass transfer coefficients, higher heat transfer coefficients improve efficiency for desiccants that have higher water uptake at lower temperatures.

7. Exergy analysis in combustion

Motivation Past studies have shown that the combustion process typically destroys about one third of the exergy of the fuel. The significance is even clearer when one notes (cf. [50]) that of the many processes occurring in the typical electricity-producing power plant, combustion of the fuel is the single
largest contributor of exergy losses in power production. It is thus of great interest to determine the magnitudes and causes of irreversibility in this process.

Past work summary The rigorous intrinsic exergy or entropy analysis, its analysis and some of the key past papers, are those considering premixed flames stabilized above a flat-flame burner \cite{51}, diffusion flames \cite{52}, droplet/spray combustion (cf. \cite{53}), as well as couple of papers that dealt only with the derivation of the equations \cite{54,55}. Almost no practical conclusions about the use of the analysis for process improvement were drawn.

Dunbar and Lior \cite{56,57}, have probably been the first to evaluate the primary causes for irreversibility, using a somewhat heuristic finite increment exergy analysis method for a simple hydrogen or methane combustor. With application to internal combustion engines in mind, Caton \cite{58} investigated the effects of temperature, pressure, and equivalence ratio for the combustion of octane-air mixtures in adiabatic, constant volume combustion systems on the exergy destruction. He performed a simple global control volume exergy analysis on the entire cylinder system, similar to the first step of the above-described analysis by Dunbar and Lior \cite{56,57}. He concluded that exergy losses decrease as the combustion temperature increases (made possible by increasing the temperature of the inflowing reactants), and as the equivalence ratio increases, and that they are fairly insensitive to the process pressure.

8. Droplet and spray combustion

8.1. Background

Based on the solution of the field equations in spherico-symmetrical coordinates, Hiwase, Datta and Som \cite{59} and Datta and Som \cite{60} have examined entropy generation and exergy destruction in droplet (the former paper) and sprays (the latter) combustion. The velocity and species concentration fields for the gas phase, and the temperature field for both the gas and for the droplet phases have been computed from the numerical solution of the equations of conservation of mass, momentum and heat, and the rate of entropy generation due to transport processes and chemical reaction in the gas phase has been determined from the generalized entropy transport equation similar to those shown by Eqs (5)–(9). They show time-integrated values of the exergy addition, irreversibility, and exergy efficiency, and found that, in a typical diffusion-controlled droplet combustion process, in which the rate of chemical reaction is much faster than the rates of diffusion of heat, mass and momentum, the entropy production rate due to the chemical reaction is lower than that due to diffusion processes. Consistent with the findings of the above-described simpler analyses, a low initial Damkohler number (as close as possible to its limiting value for initiation of ignition) and a high free stream temperature were found to cause less destruction of exergy. The field distribution of exergy destruction or entropy generation are not shown.

8.2. An example: exergy analysis of oil droplet combustion.

The droplet combustion process. Once the droplet is introduced in the hot atmosphere, its temperature starts to rise and consequently some of it begins to vaporize at its surface. The initial vaporization rate is slower because the droplet is cold, and it increases as the droplet heats up. Fuel vapor diffuses through the hot gas until the criterion for the combustion reaction is met. At this point the reaction starts
and the fuel is oxidized to form carbon dioxide and water. The combustion heat released raises the temperature of the gas and of the droplet. Fig. 4 illustrates the problem. As the droplet continues vaporizing, its radius decreases, thus originating a moving interface between the liquid droplet surface and the outer, gaseous, domain. The process will continue until all the fuel is consumed.

A single spherical droplet of heptane was considered, and a model was developed and solved for the transient mass, temperature, and velocity fields in the gas and droplet (with the temperature and concentration dependence of the properties taken into account), which are then used for the Second Law analysis of the problem. With that objective in mind, it was enough to use global kinetics for the chemical reactions, assuming full completion of the reaction. The reaction rate is given by the commonly used Arrhenius form equation

\[
\dot{\omega} = A \left( \frac{\rho_g Y_F}{MW_F} \right)^a \left( \frac{\rho_g Y_{O2}}{MW_{O2}} \right)^b \exp\left[-Ae/(RT_g)\right]
\]  

(24)

where \( Y_F \) and \( Y_{O2} \) are the mass fraction of the fuel and oxygen, respectively.

Radiative transfer was ignored, as it complicates the analysis significantly, and the literature has shown that the results are within 8% without it if no significant sooting occurs.

The finite element program FEMLAB was used for the solution, and a special computational scheme was developed to account for the diminution of the droplet size as it evaporated. Convergence and grid-independence of the numerical solution were proven, and the results were validated by comparison with those available in the literature. The computed distribution of temperature with space and time is shown in Fig. 5.

It shows clearly the ignition point (defined where \( T_g > 2000 \) K) and the flame location (denoted by the gas temperature surface maxima), as well as the temperature distribution through the domain as a function of time. It also shows that the high temperature zone is confined in a small region of \( r \),
specifically for $t > 0.432$ s and $0.002 < r < 0.09$ m, indicating the high temperature-gradients affecting the entropy generation.

Fig. 6, showing the gas temperature as a potential plot, complements Fig. 5 for easier visualization of the evolution of flame position and of the droplet moving surface.

8.3. Exergy analysis

A primary objective of this analysis is to calculate the exergy efficiency $\varepsilon$ defined by

$$\varepsilon = 1 - \frac{E_d}{E_{in}}$$

Fig. 6. The computed gas temperatures distribution as a function of time for an $n$-heptane droplet of $d_i = (1) 10^{-4}$ m, $T_{ni} = 293$ K burning in air at $T_{gi} = 1073$ K. (colour figure available online).
where $E_d$ is the exergy destruction in the defined control mass, and $E_{in}$ is the exergy inflow into it. The exergy analysis model control mass is shown in Fig. 7, and we first proceed to derive the equations needed for computing $E_{in}$. The exergy balance for the control mass (CM) over this time interval $dt$ is then given by the equation

$$E_{CM}(t + dt) - E_{CM}(t) = \left(1 - \frac{T_o}{T_g}\right)\delta Q_{CM} - (\delta W_{CM} - p_o dV_{CM}) - \delta E_d$$

(26)

where $E_{CM}(t + dt) - E_{CM}(t)$ is the overall exergy change in CM in the time interval $[t, t + dt]$. The terms on the r.h.s. of the equation represent the different contributions to this overall exergy change: $(1-(T_o/T_g))\delta Q$ is due to heat transfer between CM and the surroundings, $(\delta W - p_o dV)$ is due to the work interactions of CM with the surroundings, in which $\delta W$ is the work done or received by the system and $(p_o dV)$ the work necessary associated with changing the volume of CM, and $\delta E_d$ is the exergy destruction in CM. The exergy change in the gas phase, CV, is obtained after some steps for this essentially isobaric

Fig. 7. Sketch of the control mass (CM) used for exergy analysis for $t=t$ and $t=t+dt$, where $R(t)$ is the droplet radius at $t=t$, $R(t+dt)$ is the droplet radius at $t=t+dt$, $R_m$ is the radius of the domain extent, $h_1$: the surface between shaded region that contains $\delta m_F$ and the CV, $h_2$: the surface between CV and the external environment, $\delta m_F$ is the fuel mass contained in the internal shaded region which length is $\delta R_F$, $\delta m_{O2}$ is the oxygen contained in the external shaded region which length is $\delta R_{O2}$, $\delta Q$ is the amount of heat exchange with the surroundings, and $\delta W$ is the amount of work interaction with the surroundings. (colour figure available online).
systems, as
\[ \text{d}E_{cv} = (e_{h1} \delta m_F + e_{h2} \delta m_{O2}) - \delta E_d \]  
(27)
where the first term on the r.h.s. is the exergy input to the above-defined CV that can be expressed in integral form by the equation
\[ \dot{E}_{in} = \dot{n}_F [e_{F,ch} + e_{F,th} + \beta_{O2}(e_{(O2)ch} + e_{(O2)th})] \]  
(28)
where \( \dot{n}_F \) is the molar mass evaporation rate, \( e_{F,ch} \) and \( e_{F,th} \) are the fuel specific chemical and thermal exergy and \( e_{(O2)ch} \) and \( e_{(O2)th} \) are the oxygen specific chemical and thermal exergy, respectively.

To use Eq. (25), it now remains necessary to compute the exergy destruction rate \( \dot{E}_d \), where the specific entropy generation rate due to the heat and mass transfer and the chemical reactions, based on Eqs. (7)–(9) is,
\[ \dot{s}_p = \sum_{j=1}^{ng} \left[ \left( \rho_g D_g \frac{\partial Y_j}{\partial r} \right) \frac{R}{Y_j} \frac{\partial Y_j}{\partial r} \right] + \frac{1}{T_g} k_g \left( \frac{\partial T_g}{\partial r} \right)^2 + \frac{1}{T_g} \sum_{j=1}^{ng} s_j \left( \rho_g D_g \frac{\partial Y_j}{\partial r} \right) \frac{\partial T_g}{\partial r} + \frac{1}{T_g} \left\{ \sum_{j=1}^{ng} [h_j(T_g) - T_g s_j(T_g, P_{Vj})] \omega_j \right\} \]  
(29)
In Eq. (29) the first term accounts for mass transfer (\( \dot{s}_m \)), the second is due to heat transfer (\( \dot{s}_h \)), the third is the product of the coupling between heat and mass transfer (\( \dot{s}_c \)) and the fourth takes into account the production due to the chemical reaction (\( \dot{s}_{ch} \)).

Integrating Eq. (29) over the gas phase control volume \((R(t) < r < R_\infty)\), gives the total entropy generation rate at any instant in the CV as
\[ \dot{S}_p = 4\pi \int_{R(t)}^{R_\infty} \dot{s}_p r^2 dr \]  
(30)

The total amount of exergy addition (\( E_{in} \)) and irreversibility or exergy destruction (\( E_d \)) from the start \((t = 0)\) till any later time \( t \) are then calculated by integrating, over that period of time, Eq. (28) for \( \dot{E}_{in} \) and Eq. (30) (multiplied by \( T_o \)) for \( \dot{E}_d \).

Fig. 8. Computed history of (a) \( \dot{S}_{ch} \), (b) \( \dot{S}_{m} \), (c) \( \dot{S}_{h} \), and (d) \( \dot{S}_{c} \) and (e) \( \dot{S}_{ratio} \) for an n-heptane droplet of \( d_i = (1)10^{-4} \) m, \( T_{hi} = 293 \) K burning in air at \( T_{gi} = 1073 \) K. (colour figure available online).
To compare the magnitudes of the chemical reaction entropy generation relative to the other entropy generation terms we define and calculate the ratio

\[
\hat{S}_{\text{ratio}} = \frac{\hat{S}_{\text{ch}}}{(\hat{S}_m + \hat{S}_h + \hat{S}_c)}
\]  

Fig. 8a shows the contributions of each of the entropy terms composing Eq. (29), and Fig. 8b shows \(\hat{S}_{\text{ratio}}\), as a function of time. All the individual terms in the equations used to describe the entropy generation are presented in detail further below.

As the droplet evaporation proceeds, the total entropy generation increases due to the increase in the concentration gradients and the increasing rate of the chemical reaction in the gas phase caused by the higher fuel concentration that has been diffused from the droplet, as shown in Fig. 8a. Only \(\hat{S}_h\) decreases during the pre-combustion stage of the process [( ■) curve in Fig. 8a] due to the diminution of the heat gradient in this period resulting from the progressive rise of the droplet temperature while the gas phase temperature does not vary significantly (Figs. 5 and 6). Once combustion is attained (at \(t = 0.432\) s), the chemical reaction entropy component becomes the main contributor to the total entropy generation. The

Fig. 9. The computed (a) \(\hat{s}_{\text{ch}}\) as a function of position and time, (b) \(\hat{s}_{\text{ch}}\) evolution in time, and (c) Contour plot for \(\hat{s}_{\text{ch}}\); for an n-heptane droplet of \(d_i = (1)10^{-4}\) m, \(T_{li} = 293\) K burning in air at \(T_{gi} = 1073\) K. (colour figure available online).
significance of the dominant role of the chemical component is better seen in Fig. 8b. It shows variation of $S_{\text{ratio}}$ with respect to time. Fig. 9 shows the computed chemical entropy generation during the droplet life. As shown in Fig. 8, the chemical entropy generation dominates the irreversibility production once combustion is attained and therefore Fig. 9 gives a good indication of the evolution of the local irreversibility generation during the combustion process. The rapid increase that the chemical entropy generation $S_{\text{ch}}$ exhibits when and where the combustion is triggered is clearly seen, and explained in more detail below. Another observation is that the peak value of the chemical entropy generation follows the flame evolution shown in Figs. 5 and 6.

As described by Eq. (29), the chemical entropy generation term is a function of the gas temperature ($T_g$) and the reaction rate ($\omega$). The maximum of $S_{\text{ch}}$ is obtained at $t = 0.4356$ s, which is about 3.4 ms after the criterion to achieve ignition ($T_g > 2000$ K) is met. The maximum of $S_{\text{ch}}$ occurred at the same time that the maximum of $\omega$ is reached, but about 5.4 ms before the maximum $T_g$ is reached. This indicates that the maximum of $S_{\text{ch}}$ is more influenced by the reaction rate than by the gas temperature profile. The second most important contributor to irreversibility generation is $S_h$. From Eq. (29) we see that $S_h$ depends on $(T_g)^{-1}$, $k_g$, and $(\nabla T_g)^2$, and further analysis of these terms has shown that $k_g \nabla T_g$ has the dominant influence, interestingly in important part because of the variation of $k_g$ with temperature.

In an attempt to reduce the irreversibility due to the temperature gradients, the initial environment temperature was increased by 100 K, from 1073 to 1173 K, resulting in an increase of 3.7% in $\varepsilon$.

$s_m$ defined in Eq. (29) has the terms $s_{m1} = \sum_{j=1}^{n_g} \rho_g D_g \nabla Y_j$ that represents the entropy generation due to species diffusion, and

$$s_{m2} = \sum_{j=1}^{n_g} \frac{R}{Y_j} \nabla Y_j$$

that represents entropy generation due to concentration gradients, where the summations are for all the species present in the gas phase. The latter term was found to be $10^8$-fold larger than the former, in determining the value of $s_m$.

8.4. Exergy Input

It is of interest to now examine the exergy input ($E_{\text{in}}$) behavior and influence on $\varepsilon$. It is calculated using Eq. (28).

Examining Eq. (28) and realizing that the term $(e_F + \beta_{O_2} e_{(O_2)_B})$ is nearly constant during the process, it is obvious that $(E_{\text{in}})$ depends primarily on the fuel evaporation rate $\dot{n}_F$. First, there is a moderate increase correspondent to initial fuel evaporation prior to combustion, then a rapid increases due to strong heating resulting from the ignition, and finally a period of almost constant decrease due to the droplet shrinkage with a slope change at the end of the droplet life that is an indication of the beginning of the extinction process.

One way to consider affecting the $\varepsilon$ is by slowing down the reaction. This was attempted in the analysis by reducing the pre-exponential factor $A$ in the kinetics Eq. (24) to 1/3 of that in the base case value. It was found that the duration of the high entropy production during the combustion reaction is then shorter for the case where the reaction rate is slower, while the magnitude of the entropy production rate remains basically the same, but, importantly, the droplet life is much longer for this case, $t = 1.6605$ s, as compared to the other cases evaluated ($t = 1.233$ s for the base-case and $t = 0.9315$ s for
Consequently the cumulative effect of $\dot{E}_d$ is bigger, counteracting in this way the lower instantaneous entropy values and resulting in a lower $\varepsilon$ value.

Since the total magnitude of $\dot{S}_p$ is similar for the evaluated cases for their pre- and post-combustion periods, and taking into consideration that $E_{in}$ will always be the same at the end of the droplet life (because the fuel volume is the same), it is concluded that the duration of the droplet combustion is a key factor in the value of $\varepsilon$, and its reduction may increase $\varepsilon$. To reduce the duration of the droplet combustion, the reaction rate was accelerated by increasing the oxygen mole fraction in the air from 0.23 to 0.70, and this case was called ‘fast combustion’.

Fig. 10 shows $\dot{S}_p$ for all cases evaluated. The time-evolution of the instantaneous value of the second law efficiency $\dot{S}_p$ is similar for all the cases, but the period for the ‘fast combustion’ case is much shorter: the combustion of the droplet lasts only 0.459 s as compared with 1.233 s in the base case. The pre-combustion evaporation period for this case is only 18.3% of the droplet life. The total $\varepsilon$ was 73.2%, a 7% improvement of over the base case. The improvement in the efficiency is obtained despite the higher instantaneous entropy generation values during combustion in this case.

Table 1 summarizes the main results for the cases evaluated in this work.

<table>
<thead>
<tr>
<th>Case name</th>
<th>Conditions</th>
<th>$T_{gi}$ (K)</th>
<th>$Y_{O_2}$ (in air)</th>
<th>$A$ in $\omega$ Eq. (22)</th>
<th>Ignition delay time (s)</th>
<th>Exergy destruction $E_d$ (J)</th>
<th>2d Law Efficiency $\varepsilon$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base-case</td>
<td></td>
<td>1073</td>
<td>0.23</td>
<td>A</td>
<td>0.432</td>
<td>5.140</td>
<td>68.4</td>
</tr>
<tr>
<td>High-temperature</td>
<td></td>
<td>1173</td>
<td>0.23</td>
<td>A</td>
<td>0.092</td>
<td>4.749</td>
<td>70.9</td>
</tr>
<tr>
<td>Reduced $\omega$</td>
<td></td>
<td>1073</td>
<td>0.23</td>
<td>A/3</td>
<td>1.195</td>
<td>5.165</td>
<td>68.2</td>
</tr>
<tr>
<td>Fast- combust.</td>
<td></td>
<td>1073</td>
<td>0.70</td>
<td>A</td>
<td>0.084</td>
<td>4.409</td>
<td>73.2</td>
</tr>
</tbody>
</table>

Fig. 10. Exergy inflow ($\dot{E}_{in}$) as a function of time for an n-heptane droplet of $d_i = (1) \times 10^{-4}$ m, $T_{li} = 293$ K burning in air at $T_{gi} = 1073$ K.
9. An example: exergy analysis in a pulverized coal combustion process

In the course of research of a novel low NOx pulverized coal combustor concept where the flame is stabilized by radiation and conduction from and through the combustor walls, respectively

Fig. 11. Computed $\dot{\mathcal{S}}_p$ as a function of time for an n-heptane droplet of $d_i = (1)10^{-4}$ m, $T_{hi} = 293$ K burning in air at $T_{gi} = 1073$ K (●); $T_{gi} = 1173$ K (○); 1/3 of pre-exponential factor (A) in Arrhenius Eq. (2.2) (□); fast-combustion case (×). (colour figure available online).

Fig. 12. The computed temperature distribution in the RCSC. The flow is downward. Coal feed rate = 0.2 g/s, equilibrium air/fuel ratio = 1.10, inlet air/fuel temperature = 573 K. (colour figure available online).
(the Radiatively/Conductively-Stabilized Pulverized Coal Combustor (RCSC), Kim and Lior [61,62]), a rigorous intrinsic exergy analysis was also performed, and details can be found in [63]. The solution gives the 3-dimensional distribution of gas, particle and wall temperature, and radiation intensity, gas and particle velocity, and species concentrations. As stated earlier, this allows the determination of the spatial distribution of all of the components of exergy, using Eqs. (5)–(9).

At first, an order of magnitude analysis determined that the potential, strain, and kinetic exergy components can be ignored, leaving only the thermal and chemical components.

---

Fig. 13. The thermomechanical exergy distribution in the RCSC. Same conditions as in Fig. 12. (colour figure available online).

Fig. 14. The chemical exergy distribution in the RCSC. Same conditions as in Fig. 12. (colour figure available online).
The gas species here are CO₂, CO, NO, NH, H₂O, H₂, O₂, H, OH, N₂, HNO, O, N, HCN, NH₃. Among these species, the mole concentration of NH, H, O, OH, N is less than 0.1% of the total gases, and thus their effect on the thermomechanical and chemical exergy can be ignored.

The computed temperature field in the combustor, for the gas and the coal particles (separately) is shown in Fig. 11. The primary flame region is the meniscus-shaped area about 1/3 down from the top. Among other things, one can observe the gradual heating of the gas and the coal particles, by convection and radiation from the combustor walls, the narrowness of the flame zone (which is an objective of this combustor) and the radial temperature drop due to wall heat losses. Since the thermal exergy is one of the major exergy components, the temperature field is an important indicator of the magnitude of this exergy component Fig. 12.

The thermomechanical, chemical, and total exergy distribution fields are shown in Figs. 13, 14, and 15, respectively (in these figures the direction perpendicular to the axial flow direction of the combustor, Z/R is the radial one, r/R; the numbers 1 along these axis represent r/R=0 and Z/R=0). Some of the conclusions from observing these distributions are (i) the thermal exergy follows rather closely the temperature field, (ii) the chemical exergy starts decreasing when the volatiles start burning, and then decreases precipitously when the char starts burning, and (iii) the total exergy rises to a peak at the upstream face of the flame front where the temperature is high due to thermal feedback yet the chemical exergy has not been completely used up yet.

For the conditions of this study, about 30% of the original fuel exergy is destroyed in the combustion. Over 90% of exergy destruction takes place in the thin flame zone, and less than 10% destroyed downstream of the flame zone, the latter due to the temperature drop and heat loss due to the radiative and convective transport to the surrounding wall and exit. So far, only the effects of inlet air temperature, in the range of 573–973 K, were investigated; increasing it tended to slightly decrease the exergy efficiency, in part because of the increased heat losses.
While one may draw many conclusions from this analysis about possible ways to improve the exergetic efficiency in this combustor, a couple are already evident. One is the effectiveness of thermal feedback: the thermal exergy is raised by this feedback upstream of the flame front to about 1/3 of its maximal value. Another is the possibility of the existence of an optimal combustor exit location for the mixture, where the exergy is higher than the one shown here and yet the reactions were completed to satisfactory extent.

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