EcoAgriTourism, in the light of its multidisciplinary character, is a wide-open journal which brings together the opinions of specialists from both academic and economic environment, fostering fruitful collaborations.

The journal's structure covers all aspects of the fields approached, the focus being on original and current researches with applications in agriculture, food industry and rural tourism. Collaborators may feel free to undertake biological and technical aspects as well as aspects with social, cultural and environmental impact. Information of general interest is also welcome for the agricology-food-tourism axis.

Prof. Romulus Gruiu Ph. D.

The Journal of EcoAgriTourism aims at approaching analyses, methodologies, options and references within the journal's framework.

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RESEARCH CENTRE FOR ENGINEERING AND MANAGEMENT IN AGRICULTURE, FOOD AND TOURISM,
Transilvania University of Brasov
Bd. Eroilor nr. 29, 500036, Brasov,
Tel & Fax: 0268-415326, Tel: 0268-413000/161
e-mail: ecotec@unitbv.ro, http://www.unitbv.ro/at

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THERMODYNAMIC ANALYSIS OF HEAT EXCHANGING APPLIANCES

Shutyuk Vitaliy, Samiylene Sergiy, Vasylenko Sergiy

Abstract: The article examines the methodology of thermodynamic analysis and the optimization of heaters and heat exchangers of heat engineering complex of saccharine industry. The authors suggest noncyclic approach to the analysis of efficiency of heat exchanging apparatus, the basis of which is irrefutable fact that irreversibility as physical reason of inefficiency of technical heat engineering systems really exists. Thermodynamic analysis, which was mentioned in the article, assumes determination of measure of irreversibility of processes in the apparatus and energetic efficiency of apparatus in the whole with the help of exceptionally entropy. The measures taken to improve energetic efficiency of apparatus of saccharine factory on the example of the first group of juice heating in front of evaporator system using the given methodology, were fully analyzed.

Key words: thermodynamics, heat exchange, process, entropy, exergy of warmth.

1. INTRODUCTION

Heat exchanging apparatus of heat exchanging engineering system of saccharine industry are a part of general energy supply allocation of an enterprise as the major secondary heat and electrical energy user, which makes it difficult to analyze and optimize these apparatus. That consequently requires systematic approach with the usage of applicable methods.

It is obvious that nowadays such major characteristics as “area of thermoexchange surface” and “coefficient of efficiency” are traditionally used in saccharine industry. That is not enough, as while comparing constructionally different HEAs it makes no sense to compare relation between area of thermoexchanging surface and its characteristics. The usage of exergy method of thermodynamic analysis [3] (which is widely used while analyzing technical systems – work generators) contradicts the fundamental principles of methodology of optimization of thermoexchanging processes and systems.

The issue of choice of analysis of effectiveness of HEA was reviewed by the authors in [1,2], where the expediency of usage of non-cyclic entropy method for thermodynamical analysis and HEA optimization, as well as energetic balance method for composing energy model of HEA performance is substantiated.

Adhering to producers’ terminology, HEA of “condensate-juice” type is called as “heat exchangers”, and HEA of “steam-liquid” type – as “heaters”.

According to non-cycle entropy method technique [1, 2], integrated thermodynamic analysis assumes the determination of measure of irreversibility of processes, that occurs in HEA, the sources of which are heat exchanging at the finite variance of temperatures, the dissipation of mechanic energy of heat transfer medium currents and heat exchanging with the environment.

Fig. 1. Before the folding of entropy balance of HEA
The quantitative characteristics of irreversibility is increasing of entropy of isolated system, which determines from the entropy balance of ABC system (Fig. 1), which consists of 3 subsystems: A, B, and C (A is heating heat transfer medium subsystem, B - heat transfer medium subsystem, C – environment subsystem).

In general, entropy balance of HEA is agglomerated with the help of following simplifications:

- change of kinetic and potential energy is neglected;
- for heat transfer mediums, in which transition between preset thermodynamic states is followed by temperature changes (Fig. 2 a, 2 b), change of thermal qualities is not considerable, which allows to introduce medium thermodynamic temperature:

\[ T_m = \frac{T_{in} - T_{out}}{\ln \frac{T_{in}}{T_{out}}} \].

![Fig. 2. Change of thermodynamical states of hot and cold heat transfer media: a – in heat transfer medium, b – in heater](image)

Written form of entropy is grounded on its qualities and assumes that all its parts are absolute values; entropy can be either brought in or taken out together with the streams of substance and heat, and increase because of the irreversibility of the processes.

Entropy balance of subsystem A:

\[ m_1 s_1^{in} + \frac{E_{1d}}{T_{m_1}} = m_1 s_1^{out} + \frac{Q}{T_{m_1}} + \frac{Q_0}{T_{m_1}}; \]

Entropy balance of subsystem B:

\[ m_2 s_2^{in} + \frac{Q}{T_{m_2}} + \frac{E_{2d}}{T_{m_2}} = m_2 s_2^{out}; \]

Entropy balance of subsystem C:

\[ \Delta S_C = \frac{Q_0}{T_0}. \]

Estimating entropy additivity, in another words, \( \Delta S_{ABC} = \Delta S_A + \Delta S_B + \Delta S_C \), and the fact that AB subsystem together with C subsystem create generally isolated adiabatic system ABC (concluded from the boundary line quantities of the system), for which entropy change equals general entropy increase from irreversibility of the processes: \( \Delta S_{ABC} = \Delta S_{irrev}^{tot} \).

It can be written in the following way:
\[ \Delta S_{\text{tot}}^{\text{irrev}} = m_1 s_{11}^{\text{out}} + m_2 s_{22}^{\text{out}} - (m_1 s_{11}^{\text{in}} + m_2 s_{22}^{\text{in}}) + \frac{Q_0}{T_0}, \quad (5) \]

or

\[ \Delta S_{\text{tot}}^{\text{irrev}} = \left( \frac{Q}{T_{m_2}} - \frac{Q}{T_{m_1}} \right) + \left( \frac{Q_0}{T_0} - \frac{Q_0}{T_{m_1}} \right) + \frac{E_1^d}{T_{m_1}} + \frac{E_2^d}{T_{m_2}}. \quad (6) \]

We rewrite the equation (6) in general view:

\[ \Delta S_{\text{tot}}^{\text{irrev}} = \Delta S_{\text{tot}}^{T} + \Delta S_{\text{tot}}^{0} + \sum \Delta S_{\text{tot}}^{p}, \quad (7) \]

In which \[ \Delta S_{\text{tot}}^{T} = \frac{Q}{T_{m_2}} - \frac{Q}{T_{m_1}} \] is increasing of system entropy, conditioned by irreverence of heat exchange between subsystems A and B; W/K;

\[ \Delta S_{\text{tot}}^{0} = \frac{Q_0}{T_0} - \frac{Q_0}{T_{m_1}} \] - increasing of entropy at dissipation of mechanical energy of streams of heat transfer medium (in case of the heaters \[ E_i^d / T_{m_i} = 0 \]) W/K.

Thermodynamical efficiency of HEA, considering the irreverence of the processes, is defined by non-dimensional coefficients: entropy coefficient of thermodynamical efficiency:

\[ \eta^p_i = 1 - \frac{\Delta S_{\text{tot}}^{\text{irrev}}}{\Delta S_{\text{tot}}^{\text{max}}}; \quad (8) \]

Or entropy coefficient of thermodynamical non-efficiency:

\[ \eta^{\text{imp}}_i = \frac{\Delta S_{\text{tot}}^{\text{irrev}}}{\Delta S_{\text{tot}}^{\text{max}}}, \quad (9) \]

While

\[ \eta^{\text{imp}}_i + \eta^p_i = 1. \quad (10) \]

In which \[ \Delta S_{\text{tot}}^{\text{max}} \] is increasing of entropy of isolated system, that goes across to two given states, W/K; \[ \Delta S_{\text{tot}}^{\text{max}} \] – maximal possible increasing of entropy of adiabatic system – system passes from given state to the state of thermodynamic balance with the environment, W/K.

Coefficients (8) and (9) do not have known (discovered) drawbacks of performance factor (energetic, exergy), as they characterize the degree of diversion of real system from reverence in structure borders of the second thermodynamic law. Let us explain this.

Including the fact that the state of balance of isolated system is defined by the maximum of its entropy (consequence of the second thermodynamic law) and restrictions, imposed by the nature on the operation of technical systems (energy has technical meaning till it has the potential different from the one of the environment), denominator of equations 8 and 9 is used as the standard of comparison.

It means that \[ \Delta S_{\text{tot}}^{\text{max}} \], being the result of heat exchanging of hypothetical TS (system A) with the environment (system C) (pic. 3), which quantitatively characterizes maximal irreversibility at given characteristics of environment and is calculated with the help of the following equation (Fig. 3):
\[ \Delta S_{irrev}^{\text{max}} = m_1(s_0 - s_1) + \frac{Q_0}{T_0} + \frac{E_1^d}{T_0} + \frac{E_2^d}{T_0}, \]  

(11)

In which \(s_0\) - is specific entropy of heating heat transfer medium at the temperature of environment, J/(kg/K).

While analyzing the heaters in equation (6) 
\(T_{m1} = T_{1s}\) - the temperature of saturation of dry saturated steam, and \(\frac{E_1^d}{T_0} = 0\).

The efficiency of functioning of HEA - local effectiveness of potential usage of heat transfer medium (temperature), including dissipation processes in given temperature interval – defined entropy coefficient of HEA effectiveness:

\[ \eta_s = \frac{\Delta S_{irrev}^{\text{min}}}{\Delta S_{irrev}^{\text{tot}}} . \]  

(12)

In which \(\Delta S_{irrev}^{\text{min}}\) - is a minimal entropy increasing because of heat exchanging irreverence in HEA, 
\(\Delta S_{irrev}^{\text{min}} = Q(T_{m1}^{\text{min}} - T_{m1})/(T_{m1}^{\text{out}} - T_{m1})\), W/K, \(Q\) - true heat effectiveness of HEA, W; \(T_{m2}\) - medium thermodynamic temperature of the heating up heat transfer medium, K; \(T_{m1}^{\text{min}}\) - minimal possible medium thermodynamic temperature of heating heat transfer medium, K. For heaters \(T_{m1}^{\text{min}} = T_{1s}^{\text{min}}\) (Fig. 4 a); for heat transfer media with large mass account thermal capacity of heating heat transfer medium according to equation 6 and pic. 4 b 
\(T_{m1}^{\text{min}} = T_{2}^{\text{in}}\); for heat exchangers with larger mass account thermal capacity of heating up heat transfer medium \(T_{m1}^{\text{min}}\) is calculated analogically, including \(T_{1}^{\text{in}} = T_{2}^{\text{in}}\) (Fig. 4 c).

Fig. 3. Hypothetic TS

Fig. 4. Before defining \(\Delta S_{irrev}^{\text{min}}\): a - in the heater; b – in the heater with bigger mass consumptive thermal capacity of heat transfer medium; c – in heater with bigger mass consumptive thermal capacity of heating heat transfer medium
2. ANALYSIS OF GROUPS OF HEATERS BASED ON DEVELOPED COEFFICIENTS $\eta_{hp}$ AND $\eta_e$

Let us consider detailed features of usage of suggested coefficients with an example of heaters' analysis of the first group of the juice heaters in front of evaporator system for saccharine factory with efficiency of 6000 tons of beet processing per day with heading the standards for increasing their efficiency.

Parameters of job of heating group: juice expense – 118 % up to beetroot mass; juice temperature in front of the heater – 87 ºC; temperature of saturation of the heating steam – 107 ºC.

![Image](image_url)

Fig. 5. Relation between entropy coefficient of thermodynamic efficiency of group of heating of the heaters and area of heat exchanging surface

![Image](image_url)

Fig. 6. Relation between entropy coefficient of efficiency of heating group of the heaters and area of the heat exchanging surface
3. INCREASING OF THE AREA OF HEAT EXCHANGE \( F \) WITH THE VALEU OF \( K \).

By calculation and analysis of effectiveness of heating groups, that consist of one, two, three sequentially by juice linked ten-course heaters with \( F = 300 \) \( \text{m}^2 \) of every single one (PDS-10-300), the velocity of juice motion in pipes equals \( \omega = 1,38 \) \( \text{m/s} \) (the scum formation is not included). On picture 5 and 6 dynamics of coefficient variation with \( F \) increasing at constant \( k \) is mentioned. As it was expected, with increasing of \( F \) both the coefficients increase to some maximum value, which is caused by the defining influence on general irreverence of reduction of irreverence of heat exchange. In addition, although the range in coefficients’ change is different, maximum of the curve of each of them corresponds equal area of heat exchange.

4. INCREASING OF COEFFICIENT OF HEAT TRANSFER \( K \).

4.1. In heaters with hydraulically smooth pipes (ducts) – increasing of liquid motion velocity.

4.1.1 The effectiveness of heating groups, that consist of one, two, three sequentially switched on juice ten–coursed heaters with \( F = 200 \) \( \text{m}^2 \) for each (PDS-10-200), was calculated and analyzed. The juice velocity in pipes equals \( \omega = 2,08 \) \( \text{m/s} \). The results, given on picture 5 and 6, show that the value of coefficients are less than those received in previous case for all the range variation of \( F \), but the dynamics of their change is bigger (depends from the velocity), it is vividly demonstrated by \( \eta \) (Fig. 6).

At the same time, extremum is expressed more accurately and it is biased left.

CONCLUSIONS

Suggested technique of thermodynamic analysis assumes scientifically proved systematic approach to comparative analysis and different construction, that, obviously, is suitable to do with the help of entropy coefficient of efficiency, as well as for defining their thermodynamic efficiency in margins of sugar plant. The last can be achieved with the help of using entropy coefficient of thermodynamic efficiency and allows to analyze different heat exchanging systems for defining the level of their influence on general energetic efficiency of sugar plant.

REFERENCES

