ABSTRACT

This paper presents both criticism and suggested changes to boiler efficiency standards associated with fossil-fired steam generators. These standards include the widely used ASME PTC 4.1 and DIN 1942, and their replacements ASME PTC 4:2008 and the European EN 12952-15. For these standards it is useful to review both old and new. The chief criticism lies with inconsistent application of thermodynamic principles. Conceptual errors are made with reference temperatures and with shaft powers. This paper advocates for the Input/Loss Method.

When using computed fuel flow as a touchstone, it becomes obvious that arbitrary use of reference temperatures and/or use of capricious energy credits cannot produce a consistent (absolute) computed fuel flow. Efficiency, calorific value and fuel flow must have fixed definitions concomitant with a system’s useful energy flow. Thermodynamics is not an arbitrary discipline, the computed fuel flow of a system must describe the actual needs. Boiler efficiency requires consistent treatment, producing consistent and absolute fuel and emission flows.

Boiler efficiencies and associated calorific values have obvious standing when judging contractual obligations, for thermal performance monitoring, and for confirming carbon emissions. Note that a 0.5 to 1% change in efficiency may well have significant financial consequences when testing a new unit, or the on-going costs associated with fuel and carbon taxes. This paper demonstrates that errors greater than 2% are entirely possible if following the current standards. This paper appeals to the resolution of efficiency at the 0.1% level.

The power plant engineer is encouraged to read the Introduction and Summary & Recommendations sections while the thermodynamicist is requested to thoroughly review and critique the mid-sections. The author hopes such reviews, at a minimum, will advocate for more open discussion.

NOMENCLATURE

Note that much of the following nomenclature is taken from Exergetic Systems’ Input/Loss Method and its steam generator simulator, EX-FOSS (Lang, 2012a).

Molar Quantities Related to Stoichiometrics

- $b_A$ = Moles of water in combustion air, moles/base
- $b_Z$ = Moles of water in-leakage, moles/base
- $b_S$ = Moles of pure sorbent injected, moles/base
- $n_j$ = Moles of product j at boundary, w/o leakage
- $n_{\text{Ideal-j}}$ = Moles of ideal product j, without leakage
- $n_{S-H_2O}$ = Moles of sorbent product hydrate at boundary
- $N_k$ = Molecular weight of substance k
- $x$ = Moles of fuel/100 moles dry product (base)
- $z_S$ = Moles of H$_2$O per sorbent product
- $z_H$ = Moles of hydrogen per gaseous fuel
- $\alpha_k$ = Moles of As-Fired fuel constituent k
- $\beta$ = Molar ratio of air leakage to combustion air
- $\gamma_S$ = Moles of excess sorbent per pure sorbent $b_S$

Quantities Related to System Terms

- $CV$ = Calorific Value
- $HBC$ = Firing Correction relative to $T_{\text{CAL}}$, $\Delta$Btu/lb$_{AF}$
- $HHV$ = Fuel gross CV at constant volume, Btu/lbm$_{AF}$
- $HHVP$ = As-Fired gross CV corr. for constant pressure
- $HNSL$ = Non-Chemistry & Non-Stack Losses.
- $HPR_{\text{Act}}$ = Enthalpy of Products, actual As-Fired.
- $HPR_{\text{Ideal-XX}}$ = Enthalpy of ideal products at $T_{\text{CAL}}$, see “Subscripts” for XX, Btu/lbm$_{AF}$.
- $HRX_{\text{Act}}$ = Enthalpy of Reactants, actual As-Fired.
using an Input-Output Method is fundamentally:

statement lies with fuel water plus fuel hydrogen (thus

of a fossil fuel dictates the method by which the

means that the method of determining the energy content

more fully explained in subsequent sections, this statement

standard addresses calorimetric fundamentals. Although

INTRODUCTION TO STANDARDS

HHV = Gross calorific value, higher heating value.

CM = Calorimetric Fuel with Moist Air (and other

AF = As-Fired fuel (wet with mineral matter).

CAL = Calorimetric, as in calorimetric temperature.

CF = Calorimetric Fuel (wet with mineral matter)

 referenced to dry O\textsubscript{2} at T\textsubscript{CAL}.

CM = Calorimetric Fuel with Moist Air (and other

reactants) all at T\textsubscript{CAL}.

HHV = Gross calorific value, higher heating value.

LHV = Net calorific value, lower heating value.

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HHV = Gross calorific value, higher heating value.

LHV = Net calorific value, lower heating value.

HRX\textsubscript{CAL-XX} = Enthalpy of generic reactants at T\textsubscript{CAL}.

J = Energy conversion, 778.16926 ft-lbf/Btu

LHV = Fuel net CV at constant volume, Btu/lbm\textsubscript{AF}

LHVP = As-Fired net CV corr. for constant pressure

m\textsubscript{AF} = As-Fired fuel mass flow rate, lbm\textsubscript{AF}/hr

Q\textsubscript{WF} = “Useful Energy Flow Developed” to working

fluid from combustion gases, Btu/hr

R = Gas constant, 1545.325 ft-lbf/lbm-mole/R

T\textsubscript{CAL} = Calorimetric temperature, °F

T\textsubscript{RA} = Ambient air temp., ref. for PTC 4.1 and 4.4, °F

T\textsubscript{Stack} = Exit (Stack) boundary temperature, °F

W\textsubscript{ID} = Fan powers regards outlet streams, Btu/hr

\(\eta\textsubscript{A} = \text{Boiler absorption efficiency, unitless}\)

\(\eta\textsubscript{B} = \text{Boiler efficiency, unitless}\)

\(\Delta H\textsubscript{AF-CaSO\textsubscript{4}} = \text{Heat of Association of CaSO\textsubscript{4} hydrates}\)

\(\Delta H\textsubscript{AF-CaSO\textsubscript{4}} = \text{Heat of Disassociation (e.g., Trona) at T\textsubscript{CAL,}} \Delta Btu/lb-mole.\)

\(\Delta H\textsubscript{CAL} = \text{Heat of Form. of k at T\textsubscript{CAL,} \Delta Btu/lb-mole}\)

\(\Delta H\textsubscript{CAL-H2O} = \text{Heat of Formation, sat. liquid at T\textsubscript{CAL}}\)

\(\Delta H\textsubscript{CAL-H2O} = \text{Heat of Formation, sat. vapor at T\textsubscript{CAL}}\)

\(\Delta H\textsubscript{CAL} = \text{Enthalpy correction for net CV,} \Delta Btu/lbm\textsubscript{AF}\)

\(\Delta H\textsubscript{PV} = \text{Enthalpy correction for volume,} \Delta Btu/lbm\textsubscript{AF}\)

\(\Delta P\textsubscript{L/H} = \text{PV energy correction from net to gross CV}\)

\(\Delta U\textsubscript{L/H} = \text{Internal energy correction from net to gross.}\)

The boundary of a fossil-fired steam generator being studied for thermal efficiency does not encompass all physical equipment, but rather the constraining volume of its interacting fluids, for example: fuel conveyance, combustion gas confinement, the inside of air ducts and working fluid pipe IDs. This boundary derives from the principle that the thermal efficiency of a steam generator only addresses how the As-Fired fuel interacts with its gas/air/working fluids.

Extraneous equipment has no impact on boiler efficiency if not directly affecting the fuel’s interaction with the gas/air/working fluids. This definition is consistent with the concept of the fuel’s calorific value. It immediately excludes such equipment as pulverizers, steam driven pumps, recirculation pumps, and the like. ASME PTC 4 and DIN 1942 would suggest that a higher pulverizer shaft power will increase boiler efficiency. However, pulverizer power has no impact on the interaction of fuel and gas/air/working fluids (the grinding of coal is emulated during lab preparation of samples). A recirculation pump has no impact on fuel heating the working fluid. The metric boiler efficiency must solely guide the engineer towards reduced fuel flow and CO\textsubscript{2} emissions while making adequate steam. Shaft powers are monitored through house load.

A system in equilibrium with its medium, from which no power is extracted, defines its dead state (Keenan, 1941). For an active steam generator, having the potential for thermal power, the condition of its dead state is often confused with its “reference state”. The dead state limits how much potential power is possible, an ultimate limit to actual output, the numerator of Eq.(1). For steam generators, the dead state should be taken as the coldest medium associated with the local environment. For an actual steam generator, its thermal power as derived from a given fuel flow are absolutes, established by considering unambiguous differences between reference states (main steam less feedwater, etc.). In engineering, all energy levels are relative to a chosen reference state. Text books will argue that reference states are arbitrary, a simple enough concept when dealing with a single fluid. Water
properties for example, normally referenced to the triple point, could be referenced to the boiling point at 1 atmosphere, resulting in the same Useful Energy Flow Developed of Eq.(1).

However, for this discussion we are not concerned with the numerator of Eq.(1), but rather treatment of the denominator. Note that although Eq.(1) expresses an Input-Output approach, conversion to specific values immediately invokes a Heat Loss Method (herein termed the “Energy Balance Method”); but the two must produce identical efficiencies. The argument here is for absolutes in the denominator, that the “arbitrariness” of reference states can not be defined by the casual analyst but, indeed, is established when defining the energy content of the fuel. The technician determining calorific value of natural gas may be required run his/her calculations using a national standard, e.g., 0.0°C for France, 15°C for Ireland and the U.K., 25°C for Germany, 60°F (15.56°C) for North America, etc. (see ISO 12213-3:2006, ISO 13443:1996 /Cor 1:1997, and AGA Report No. 3), or another yet. For solid and liquid fossil fuels the reference temperature may be viewed by some as arbitrary, but in fact is also set by the technician running the bomb calorimeter, not the casual analyst. For an adiabatic or isoperibol bomb calorimeter, its reference is the “calorimetric temperature” at which the bomb’s water jacket is kept in an equilibrium state.

It is well understood that calorific values vary with temperature. Natural gas evaluated at 0.0°C is different than that at 25°C, a bomb run at 50°F will yield a different value for the same coal as one run at 120°F. Because As-Fired conditions are relative to a reference temperature, boiler efficiency will vary with reference temperature. That said, the objective of this paper - and the proposed objective of all efficiency standards - is an absolute understanding of a steam generator’s thermal performance, the fuel it consumes, consistent with a defined boundary and actual emission flow. If Eq.(1)’s numerator is an absolute measure of thermal power, we can not allow an arbitrary denominator (implying an arbitrary fuel flow). However, recognizing that boiler efficiency varies with temperature is no justification for standards advocating relative efficiencies, but rather, efficiency equations which properly correct for reference temperature. The test for absolutes is computed fuel flow, a fuel mass flow which has been corrected to the actual firing conditions and thus producing consistent, absolute, fuel & emission flows.

The act of choosing a calorimetric temperature to either compute (for gaseous fuels) or to measure (for solid and liquid fuels) a calorific value does not give the casual analyst the liberty to then choose another for reference. For any multi-fluid system, only a consistent thermodynamic reference state can be considered or the laws of thermodynamics will not be satisfied. It is not acceptable to assume one energy level for the fuel, another for the working fluid, another for the combustion products; sorbents, tube leaks and soot blowing add to products; the air’s moisture affects products; and combustion products heat working fluid.

ERRORS, WHAT ERRORS?

Addressed are three types of errors and uncertainties involving thermodynamic concepts, procedural problems, and measurements which impact efficiency:

1) System conceptual errors are made when:
   a) mis-using reference temperatures;
   b) when treating certain system components, especially shaft powers (i.e., “credit” terms); and
   c) when not understanding the energy of reactants.

2) Procedural errors are made which impact thermodynamics in a generic manner; e.g., the definition of dry air, use of consistent molecular weights, enthalpic values of fluids which are mixed within the system, thermodynamic properties, etc.

3) Measurement uncertainties such as those found in the lab regards fuel chemistry and CVs, certain loss terms (HNSL), and uncertainties in plant data which affect QWF and thus back-calculated fuel and emission flows.

For all of these, the author argues for an accuracy criteria at the ±0.1% ΔηB level. What this means is that system concepts and procedures affecting a computational efficiency at ≥ ±0.1% ΔηB must be included. This is not to say all system concepts and procedures are to be understood at this level, but are to be included given a computational sensitivity thus demonstrated. Although coal analysis between laboratories should lie within ±100 ΔBtu/lbm (±233 ΔkJ/kg), modern bomb calorimeters are quoted as having 0.1% HHV/HHV repeatability. It is this fact which adds imperative. The author’s EX-FOSS program allows for an error calculation based on stoichiometric inconsistencies (as-tested emissions versus assumed fuel chemistry), which if > ±0.1% ΔηB, and unexplained, results in repeating the test. It is noteworthy that ASME PTC 4 (§5-7.3) expresses this same opinion, that “a convergence limit of 0.1% efficiency is sufficient”.

A clear example of the misuse and capricious nature of current boiler efficiency standards is witnessed by the comments made at a 2009 TAPPI conference. The conference authors appear to be searching for a standard which would produce the highest efficiency to minimize “taxation and legislative problems”; their comments are stunning (Vakkilainen & Ahtila, 2009).
Set out below are errors made in traditional boiler efficiency standards. This list is by no means complete, but concentrates on “credits” and reference temperatures.

**ASME PTC 4.1 (United States):**
This code was superseded in 1998 by ASME PTC 4. Although PTC 4.1 is no longer an ASME code nor an ANSI standard, it is included here given its continued use and historic precedence.

§5-19.8, the enthalpy of natural gas should be computed, §5-15.5.1 and §5-15.5.2, shaft powers are included as energy credits.

§7.2, the reference temperature is taken as the air’s ambient temperature ($T_{RA}$).

§7.2.8.1 and Appendix, nitrogen content in dry air is fixed at 76.85% (an assumed weight fraction).

§7.3, “heat credits” appear in the numerator and denominator.

The FD and ID fans are considered outside the thermodynamic envelope.

The standard’s Input-Output and Heat Loss Methods are inconsistent (see the DIN 1942 discussion).

This standard applies to any fossil fuel.

**ASME PTC 4 (United States):**
The original 1998 release was superseded with a 2008 release to which the following are referenced.

§5-5, the defined fuel energy flow ignores the as-fired state (if different from 77°F), this would especially destroy any accuracy of gas-fired efficiencies given typical firing from 45°F to 60°F.

§5-7, “fuel efficiency” is PTC 4’s preferred method, fuel energy is not corrected to the as-fired.

§5-7.1, “credits” appear only in the numerator.

§5-8.1, the conversion from constant volume to constant pressure ignores nitrogen and oxygen bound in the fuel, oxygen is an important term when considering high oxygenated fuels (such as PRB); such PTC 4 conversion is not temperature dependent.

§5-9, sorbent energy flows are normalized to the as-fired fuel, which ignores unique sensible energies.

§5-9.5.1, §5-9.5.2, §11.1, §5-11.4.1, etc., oxygen content in dry air is fixed at 20.95% (Ar and CO$_2$ are not included, molar air/oxygen = 3.7733).

§5-11.2, psychrometric properties should be extended to -40F, as applicable for steam generators found in norther climes; caution should be exercised when using reference psychrometric temperatures; the standard employs a water to dry air ratio of 0.6220 (i.e., textbook), PTC 4 data would suggest 0.6398.

§5-13.1, the reference temperature is set constant at 25°C.

§5-15.5.1 and §5-15.5.2, shaft powers are included as energy credits.

§5-19.8, the enthalpy of natural gas should be computed, and is easily done given known properties.

§5-19.11 and §5-19.12, “average” combustion gas properties are erroneously employed, such use is hardly justified given the wide applicability this standard is assumed to have; the referenced 1971 JANAF/NASA properties are out-of-date (other citations, appearing political, are quite current).

The FD and ID fans are considered outside the thermodynamic envelope.

The 2008 release (unlike the 1998 version) relates to coal-, oil- and gas-fired steam generators.

**DIN 1942 (German):**
This code was superseded by DIN EN 12952-15:2004.

The following nomenclature and comments are specific to DIN 1942 (Feb. 1994).

§6.2, the reference temperature ($t_o$) is set at 25°C. However, “other temperatures may be agreed upon” by correcting the net calorific value with fuel, air and combustion gas sensible heat terms.

§6.3.2.3, so-called “heat credits” (denoted as $Q_Z$) includes shaft powers from pulverizers, recirculating gas fans, working fluid circulating pumps and “power from any other motors”.

§6.4, DIN 1942 employs the $Q_Z$ term in both its Input-Output and Heat Loss Methods. In DIN 1942: $Q_N$ is the useful output (herein $Q_{WF}$); $Q_{ZB}$ is the fuel energy ($m_{AF}LHV$); and $Q_{VTot}$ is the loss term.

\[
\eta_{B-LHV} = \frac{Q_N}{Q_{ZB} + Q_{Z}} \quad (DIN-144)
\]

\[
\eta_{B-LHV} = 1.0 - \frac{Q_{VTot}}{Q_{ZB} + Q_{Z}} \quad (DIN-147)
\]

For Eq.(DIN-147) an increase in $Q_Z$ will always increase $\eta_{B-LHV}$ provided $Q_{VTot}$ > 0. However, the same increase in $Q_Z$ will always decrease the Input-Output efficiency of Eq.(DIN-144), thus guaranteeing inconsistent computed fuel flows.

This same conundrum exists with PTC 4.1, with PTC 4 and its “gross efficiency” definition, and with other standards.

§6.3.4.1, oxygen content in dry air is fixed at 20.938% (Ar is not included, air/oxygen = 3.7760).

**Draft European Standard:**
The following comments reference prEN 12952-15 of Nov. 1999, which is now issued as EN 12952-15.
The draft and the new closely follow DIN 1942, employing its nomenclature and general methods.

§7.2, the reference temperature ($t_o$) is set at 25°C, but “other temperatures may be agreed upon” which corrects heat credits as done in DIN 1942.

§7.3.4.1, oxygen content in dry air is fixed at 20.938% (Ar is not included but apparently CO$_2$ is included,
For recovery boilers, burning black liquor fuel, is it §7.1.4.1 and §7.1.4.3, oxygen content in dry air is fixed (0) (page 4), the reference temperature is set at 77°F. This standard was prepared by the Technical Association for Recovery Boilers Used in the Pulp & Paper Industry (United States): This standard applies to any fossil fuel.

BS 2885 (British):
The following comments reference BS 2885:1974; which has been superseded by the British Standards Institute (BSI) issue of BS EN 12952-15:2003 (basically the revised DIN 1942). §2 (bottom), all fuels shall use a “calorimetric temperature” of 25°C. Section E and Section F, Item 434, the standard expects that flue gas nitrogen is to be measured, thus dry air is not specified.

Other standards reference BS 2885 (British): BS EN 12952-15:2003 (bottom), all fuels shall use a “calorimetric temperature” of 25°C. Section E and Section F, Item 434, the standard expects that flue gas nitrogen is to be measured, thus dry air is not specified.

Items 708, 804 and 907 (and Notes), the reference temperature for sensible heats in the dry flue gas, moisture in the combustion air and fuel is the combustion air temperature (per ASME PTC 4.1), not 25°C as its stated “calorimetric temperature”. Item 901 (and Notes) regards “Method A” (Input-Output) it does not consider energy credits; it invokes a simple “fuel efficiency” in which the CV bears no Firing Correction. However Item 902, invoking the Heat Loss Method (“Method B”) considers the “heat equivalent of auxiliary power” as a loss, carrying the same sign as the radiation & convection loss. Such inconsistencies will result in impossible differences in computed fuel flows.

Energy credits (shaft powers) appear only in the denominator of its Heat Loss Method. This standard applies to any fossil fuel.

This standard was prepared by the Technical Association of the Pulp and Paper Industry (TAPPI,1996) and was based on a 1993 draft of ASME PTC 4 using its energy balance method; however there appears to be some controversy in Europe when applying this standard (Vakkilainen & Ahtila,2009). §0 (page 4), the reference temperature is set at 77°F. §7.1.4.1 and §7.1.4.3, oxygen content in dry air is fixed at 23.14% (a carry-over from ASME PTC 4.1, but used as a molar ratio and is not correct).

For recovery boilers, burning black liquor fuel, is it common industrial practice to correct the measured heating value for Heats of Formation for the reduction of Na2SO4. Such corrections address the difference between ideal combustion products associated with a bomb calorimeter versus actual products associated with further reduction of certain black liquor compounds. Such corrections are thermodynamically inconsistent, as the calorific value is corrected with a computed ΔHR term: (HHVP - ΔHR + HBC). This standard applies only to recovery boilers burning sodium-laced black liquor.

CONCEPTUAL AND PROCEDURAL ERRORS

Standards have no monopoly on conceptual errors. Previous versions of the Input/Loss Method did not properly recognize energies of reactants; it used a complicated set of corrections which now appear frenetic. The present approach (this Rev. 30) greatly simplifies when developing Eqs.(3) thru (4C). They demonstrate that gross and net reactant and product terms are identical.

Most observed conceptual errors are associated with product water, formed from the fuel’s entrained water and from bound hydrogen. Thus, conceptual errors associated with the calorimetric temperature for highly energetic fuels, with low product water, are slight and typically do not meet the 0.1% ΔηB criteria. However, conceptual errors associated with fuels producing 10% or more product water (such as high volatile B bituminous (hvBb), to PRB coal, to the lignites and peat) are appreciable, ranging from 0.2% to the 0.8% ΔηB level assuming a 18°F (10 ΔC) change in calorimetric temperature. Such errors derive from assuming a reference temperature for boiler efficiency calculations, while the calorific value was determined at another. For example, a 0.5% ΔηB error is made for Powder River Basin (PRB) coal when 25°C is the assumed reference, while the CV was determined at 35°C.

Conceptual errors may also exist when not recognizing the sensitivity of the ratio of ambient oxygen to dry air. A change of ambient oxygen from 21% to 20.5% represents a decrease of 0.253% ΔHBP, for a typical PRB fired unit assuming a constant Stack O2: a 1.128% ΔηB increase for 19% ambient. Of the steam generators tested by the author, typically 1 in 10 were found to have degraded ambient oxygen levels typically caused either by a weather inversion, still air or flue gas leakage into FD Fans. The NASA (1976) standard ambient oxygen is 20.9476% at sea level.

Conceptual errors are also made when following the current standards (low water fuels aside) which involve the treatment of shaft powers, discussed in the next section.

Procedural errors are made by not adhering to the latest thermodynamic standards. The author finds it rare that any two standards use the same molecular weights. Inconsistency is present in air psychrometric properties (see ASHRAE procedures, discussed below). One would expect to use the same fundamental methods when evaluating a combustion turbine versus a conventional steam generator (ASME PTC 22 assumes 60°F as a base, conversion is allowed per its §4.12, versus ASME PTC 4).

Precise methods offer little comfort if laboratories...
cannot record CVs with at least repeatability, if not also with accuracy. Although the 0.1% ΔT B criteria is meet in repeatability when using the modern bomb calorimeter, we must remember that variability found in CVs (say from grab sampling) may, indeed, be quite real. Standards must present a practical statistical treatment of multiple lab chemistries and CVs, as would be associated with testing a coal-fired unit. It is noteworthy that ASME PTC 4 devotes its Section 7 to uncertainty analysis. Although Section 7 is clearly amenable for academic pursuits, it is not something most power plant engineers are going to place under their pillows. The coal-fired industry needs procedures, fully integrated within the standards for evaluating a test:

1) Consistency of individual as-tested fuel chemistries, rejecting any CV given an outlying chemistry [one solution is to apply techniques afforded in the Oxy-Hydrocarbon model (see Lang & Canning, 2007)];

2) Define a steady state period by examining time plots of feedwater and fuel flows (over at least 15 min.), the averaged data resulting in QWF [although such plots have been used by the author and his colleagues for years, the precedence for this is 125 years old! (Kent, 1884)];

3) Tolerance on a proper Energy Balance Method efficiency, as based on items only affecting calorific value (i.e., sampling) and QWF; and

4) A consistent boiler efficiency, allowing the calculation of an absolute fuel flow.

CONSISTENT BOILER EFFICIENCY

The temperature used to operate a bomb calorimeter, or to compute a gaseous CV, is the beginning point for developing consistent thermodynamics. Calorific values for solid or liquid fuels are obtained either by adiabatic or isoperibol bomb calorimetry following ASTM D5865 or ISO 1928:1995(E). An adiabatic bomb calorimeter detects the gross energy liberated from ideal combustion, burned in pure O₂, by maintaining a constant water bath temperature about the bomb, which defines the calorimetric temperature, TCAL. An isoperibol bomb calorimeter detects the net energy liberated by accurately monitoring the water bath temperature, its resultant average value being TCAL. Many modern bomb calorimeters are automated to run at a programmable TCAL. The author has found various labs in North America and Europe using 27°C (80.6°F), 28.5°C (83.3°F), 30°C (86°F) and, commonly, 35°C (95°F). Up thru 2007 the author could not find any laboratory in North America or in Europe determining coal CVs at 25°C, the reference for most standards!

Boiler efficiency should be a simple reflection of what the technician, determining CV by either calculation or measurement, has produced. The calorimetric process begins with reactants, the combustion event, and ends with ideal products of CO₂, SO₂ and H₂O. This process is path dependent, the traditional path is to maintain an essentially constant temperature of the calorimeter’s heat sink. When mimicking this calorimetric process when applied to a steam generator, the thermodynamicist need only account for: losses associated with actual product streams; all reactants (moist air, sorbents, leakages, etc.); and sensible heats accounting for the fact that reactants may not be fired at the calorimetric temperature. This then is the conversion efficiency of burning fuel, delivering a “Useful Energy Flow Developed” from combustion; i.e., its interaction with gas/air/working fluid. If consistent fuel and emission flows are to be computed from boiler efficiency, QWF must only reflect heating from combustion gases. The execution of these concepts is a bit more involved.

The definition of gross calorific value (higher heating value) as based on a bomb calorimeter is the energy liberated from products formed relative to the calorimetric temperature, this includes, of course, the water produced as reduced to the liquid state. We do not measure net values (lower heating values). The internal energy liberated from a constant volume bomb is relative to the equilibrium temperature at which the bomb functioned, as described by the following:

\[ \int \partial Q_{T-Cal} = - HHV = - HHVP + \Delta H_{VP} \quad (2) \]

Note that Eq.(2) is path dependent, for a traditional bomb calorimeter, industrial practice sets this path as one having a constant bath temperature. By correcting for PV work, via \( \Delta H_{VP} \), conversion is made from a constant volume internal energy (HHV) to an enthalpy base (HHVP).

A fuel’s calorific value, after conversion, is the difference between the enthalpy of ideal combustion products \( \text{HPR}_{\text{ideal-CF-HHV}} \) and the enthalpy of the reactants \( \text{HRX} \) as ideally oxidized in bone-dry O₂, and both evaluated at that temperature at which these quantities were formed, at TCAL. From First Law conservation, Eq.(2) results in the following expressions, descriptive of a “calorimetric system” (i.e., measured or computed). The net CV base of Eq.(3B) is justified below.

\[ \text{HHVP} = - \text{HPR}_{\text{ideal-CF-HHV}} + \text{HRX}_{\text{CAL-CF-HHV}} \quad (3A) \]
\[ \text{LHVP} = - \text{HPR}_{\text{ideal-CF-LHV}} + \text{HRX}_{\text{CAL-CF-LHV}} \quad (3B) \]

However, there is additional complexity. When we either measure or compute a CV, we employ dry O₂ to produce idea products, thus there are no compound formations, reactants or products, not directly associated with the pure fuel; i.e., a calorimetric system. Of course, since all streams are at TCAL there is no sensible heat. In a calorimetric system, there is no product water formed which does not derive from the fuel; it being condensed (a
gross CV), or not (a net CV). However, when analyzing a system using moist combustion air, the Heat of Formation of the air’s water must be accounted when evaluating the actual Heat of Reactants. We must track separately that water associated with the fuel, versus ambient moisture and all other reactants. For example, consider 1.0 mole of moist methane being burnt in moist oxygen. The following suggests how \( HRX_{CAL-CF} \) must be computed:

For a gross CV base, all at \( T_{CAL} \):
\[
\{0.9[CH_4] + 0.1[H_2O]\}_{\text{As-Fired}} + 1.8[O_2] + 0.3[H_2O]_{\text{Vap}} \\
=> 0.9[CO_2] + 1.9[H_2O]_{\text{Eq}} + 0.3[H_2O]_{\text{Vap}}
\]

For a net CV base, all at \( T_{CAL} \):
\[
\{0.9[CH_4] + 0.1[H_2O]\}_{\text{As-Fired}} + 1.8[O_2] + 0.3[H_2O]_{\text{Vap}} \\
=> 0.9[CO_2] + 2.2[H_2O]_{\text{Vap}}
\]

As an example of the molar nomenclature used below, in these reactions: \( x = 1.0; \alpha_{CO_2} = 0.9; b_A = 0.3; n_{CO_2} = 0.9; \) etc. In all systems, the gross CV reflects the condensation of product water derived only from the As-Fired fuel (at \( T_{CAL} \)). Also, as a fine point, it is not credible to suggest that, for example, if a gaseous fuel contains CO\(_2\) or water, or coal contains water with a leakage of working fluid, or combustion air bears moisture, that ideal combustion products cannot be defined consistent with Eq.(3).

Thus by adding the Heats of Formation of non-fuel reactants to \( HRX_{CAL-CF} \) of Eq.(3), we form an energy of combined fuel and non-fuel reactants, all at \( T_{CAL} \), termed \( HRX_{CAL-CM} \). To balance Eq.(3), such treatment implies that the energy of ideal combustion products now reflects non-fuel reactants, termed \( HPR_{Ideal-CM} \) thus:

\[
HHVP = -HPR_{Ideal-CM-HHV} + HRX_{CAL-CM-HHV} \quad (4A1)
\]
\[
LHVP = -HPR_{Ideal-CM-LHV} + HRX_{CAL-CM-LHV} \quad (4A2)
\]

where:

\[
HRX_{CAL-CM-HHV} = HRX_{CAL-CF-HHV} + [\text{Non-Fuel } \Delta H^0_{\text{F-CAL-k}}] \quad (4B1)
\]
\[
= HHVP + HPR_{Ideal-CM-HHV} + [\text{Non-Fuel } \Delta H^0_{\text{F-CAL-k}}] \quad (4B2)
\]
\[
HHVP = [H_{\text{Ideal-co}} \Delta H^0_{\text{F-CAL-CO}} + (n_{\text{ideal-soz}} \Delta H^0_{\text{F-CAL-soz}} + (n_{\text{ideal-H2O}} \Delta H^0_{\text{F-CAL-H2O}} + b_A(1.0 + \beta)\Delta H^0_{\text{F-CAL-H2O}} + b_Z(1.0 + \gamma_S)(\Delta H^0_{\text{F-CAL-Sorb}} - \Delta H_{\text{D-CAL-Sorb}})]/(xN_{AF}) \quad (4B3)
\]

\[
HRX_{CAL-CM-LHV} = HRX_{CAL-CF-LHV} + [\text{Non-Fuel } \Delta H^0_{\text{F-CAL-k}}] \quad (4C1)
\]
\[
= LHVP + HPR_{Ideal-CF-LHV} + [\text{Non-Fuel } \Delta H^0_{\text{F-CAL-k}}] \quad (4C2)
\]

As observed, Eq.(3) is substituted into Eq.(4A) for \( HRX_{CAL-CF} \) allowing \( HRX_{CAL-CM} \) to be computed using Eq.(4B3) or Eq.(4C3). In summary, since the thermal efficiency of an ideal system described by Eqs.(3) or (4A) is unity, it follows that either Eqs.(3) or (4A) must serve as the basis for all boiler efficiency standards.

**FIRING CORRECTIONS**

When developing an expression for boiler efficiency, the As-Fired fuel’s energy content must be corrected for sensible heat relative to \( T_{CAL} \). Note that firing fuel at 10\(^{\circ}\)C or 100\(^{\circ}\)C cannot affect its interaction with the gas/air/working fluids, if properly referenced to \( T_{CAL} \). As analysts we should be able to run calorimeters at any temperature, fire the fuel at any other temperature, without bias to the concept of ideal combustion, without affecting thermodynamic principles, and still compute an absolute fuel flow. Thus by simply adding a “Firing Correction” term (HBC) to each side of Eq.(4A), we bring the reactants term to the As-Fired, maintaining unity \( \eta_B \):

\[
HHVP + HBC = -HPR_{Ideal-CM-HHV} + HRX_{CAL-CM-HHV} + HBC \quad (5A)
\]
\[
LHVP + HBC = -HPR_{Ideal-CM-LHV} + HRX_{CAL-CM-LHV} + HBC \quad (5B)
\]

Note that the signs associated with Eqs.(3), (4) & (5) yield to the convention of a positive calorific value (note that the numeric value of \( HPR_{Ideal} \) is always < 0.0, and \( HRX_{CAL} \) is typically < 0.0). The efficiency of a system described by Eq.(5) is unity. Eq.(5) is interesting in that no condition of any reactant stream will cause departure from unity efficiency given all are at \( T_{CAL} \). Bone dry or fogged combustion air, nor size of FD Fan, nor steam-air heating, nor sorbent flow, nor water leakage, etc. will affect boiler efficiency per se. Such situations only affect inlet streams. Although entering the system not at \( T_{CAL} \) and so corrected by HBC, under ideal conditions all products exit at \( T_{CAL} \).

The next developmental step is to degrade from the ideal by accounting for losses. How thermodynamic losses are grouped may be treated in any number of ways. For this work, they are based on specific energy terms (Btu/lbm), and are described by \( \Sigma \)Losses/m\(_{AF} \). If just subtracting losses from \( HPR_{Ideal-CM} \), Eq.(5) becomes unbalanced without an efficiency term. Thus use of either gross efficiency (\( \eta_{B-HHV} \)) or net efficiency (\( \eta_{B-LHV} \)) to achieve conservation. The \( HPR_{Ideal-CM} \) term being at \( T_{CAL} \).
which does not maintain this concept will produce error.

Any consideration of a Firing Correction Corrections can not affect departure from unity efficiency if they correct to the As-Fired condition. Since Firing LHVP + HBC - HPR

was substituted for [HRX

Ideal-CM-LHV]

CAL-CM-LHV

HHVP + HBC

The computed boiler efficiencies follow directly, noting that in Eq.(7) the actual Enthalpy of Reactants (HRXAct) was substituted for [HRXCAL-CM + HBC]:

\[
\eta_{B-HHV} = \frac{-\text{HPR}_{\text{Ideal-CM- HHHV}} - \sum \text{Losses/m}_\text{AF} + \text{HRX}_{\text{Act-CM-HHV}}}{\text{HHVP} + \text{HBC}}
\] (7A)

\[
\eta_{B-LHV} = \frac{-\text{HPR}_{\text{Ideal-CM-LHV}} - \sum \text{Losses/m}_\text{AF} + \text{HRX}_{\text{Act-LHV}}}{\text{LHVP} + \text{HBC}}
\] (7B)

In summary, Firing Corrections are to be judged only if they correct to the As-Fired condition. Since Firing Corrections can not affect departure from unity efficiency (only losses), any consideration of a Firing Correction which does not maintain this concept will produce error.

Any standard not specifying the use of a Firing Correction term, based on calorimetrics, applied in both numerator and denominator of a proper Energy Balance equation, per Eq.(7), is in error. The concept of “fuel efficiency” is meaningless for industrial standards. It is remarkable that as reaffirmed in 1991, ASME PTC 4.1 employed “heat credits” in both its numerator and denominator, and not seven years later PTC 4 employed “credits” only in the numerator.

A standard allowing for pulverizer electrical power to be added as a “credit” is an error. Pulverizer shaft power adds nothing to the process of fuel interacting with gas/air/working fluid, other than preparing coal’s surface energy for combustion. However this action is replicated when preparing a sample for bomb calorimetry. Grinding a brittle substance affects its surface energy, not internal energy; if not ground to a optimum surface/mass, its full chemical energy cannot be realized. If HHV differences between mesh sizes used in the lab versus that produced by pulverizing could be determined, then a correction to HHV (not the Firing Corrections) is entirely justified. Although such direct comparisons are quite difficult as coal’s mineral matter grinds differentially, work was initiated which may be helpful in developing a corrective chart for HHV if mesh sizes are greatly different (Lang, 2011 and its Figs. 3 & 4).

A standard allowing the ambient air temperature to be taken as the reference is in error. For example, some standards would allow a natural gas CV to be computed at 60°F, referenced at 95°F and fired at 45°F with a computed fuel flow known but to God. The only applicable reference temperature for such a case is the calorimetric at 60°F.

A standard not appreciating that the reference states of the various fluids (working fluid leakage, gas, dry air, air’s moisture, etc.) must be referenced to the same condition, is allowing for First Law violation; steam generators mix all fluids. For example, North America’s ASHRAE psychrometric properties share a unique strangeness, where dry air is taken from 0.0°F (-17.8°C) and water from its triple point; this is absurd when addressing boiler in-leakage. For more strangeness, ISO 12213, reflecting national standards, specifies that different reference temperatures may be employed for natural gas CV versus that used for volume metering (for example, Germany employs CVs at 25°C, but flows are metered at a 0.0°C reference).

A boiler standard which allows, for example, circulating pump power to affect boiler efficiency is in error. A working fluid pump power associated with the steam generator must be subtracted from QWF, as QWF must only reflect the Useful Energy Flow Developed to the working fluid from combustion gases. Such manipulations should be kept apart from a determination of efficiency.

BOILER EFFICIENCY

The individual terms in these expressions may now be defined and expanded; see Lang (2009a, 2009b) for details. It is tacitly assumed that back-correcting an as-measured calorific value (HHV) of a solid or liquid fossil fuel to some chosen Tcal is not possible given chemical complexities of such fuels. It is for this reason that the reactant term \( \text{HRX}_{\text{CAL-CF}} \) is evaluated using Eq.(3).

As regarding loss terms, although differing techniques may be employed, the above fundamentals must be followed; specifically anything affecting an ideal product stream (i.e., \( \text{HPR}_{\text{Ideal-CM}} \)) is a loss and will decrease efficiency and increase computed fuel flow. Different procedures divide the \( \sum \text{Losses} \) term of Eq.(7) into different categories. Exergetic Systems separates \( \sum \text{Losses} \) into effluent (stack) related quantities, reflected in its \( \text{HPR}_{\text{Act}} \) term, and non-effluent quantities termed “Non-Chemistry & Non-Stack Losses” (HNSL).
\[
\eta_{B-hHV} = - \frac{\text{HPR}_{\text{Act-HHV}} - \text{HNSL} + \text{HRX}_{\text{Act-HHV}}}{\text{HHVP} + \text{HBC}} \quad (11A)
\]

\[
\eta_{B-LHV} = - \frac{\text{HPR}_{\text{Act-LHV}} - \text{HNSL} + \text{HRX}_{\text{Act-LHV}}}{\text{LHVP} + \text{HBC}} \quad (11B)
\]

This separate treatment allows definition of the actual product term without differentiating “stack losses” (as used in the standards) from actual combustion products; it eliminates formulation problems of PTC 4, EN 12952-15, DIN 1942, etc. The actual quantities are evaluated using Eqs.(13) and (14) in which standard Heats of Formation are converted to \(\Delta H_{f,CAL}^0\), referenced to \(T_{\text{CAL}}\). Conventional “stack losses” may be back-calculated for user convenience. \(\text{HPR}_k\) is the enthalpy of a non-water product \(k\) at the boundary, including sorbent products for example. Eq.(13) includes water terms, again specified separately for clarity. In Eq.(13): the term \(n_{\text{H}_2\text{O}}\) describes all water found at the boundary, less moisture carried with air leakage \((b_\text{A} \beta)\); \(b_\text{A}\) is moisture carried with combustion air (relative to \(h_{\text{g-CAL}}\)); \(b_z\) describes working fluid leakage air (relative to \(h_{\text{g-CAL}}\)); and by example, the term \(n_{\text{H}_2\text{O}} z_{\phi}\) describes water products from sorbents (e.g., \(\text{CaSO}_4\) hydrates) with a required Heat of Association given Eq.(14) would address \(\text{HPR}_{\text{CaSO}_4}\). Eq.(12) is not used, it is presented for clarity:

\[
\text{HPR}_{\text{Act-HHV}} = \text{HPR}_{\text{Ideal-CM-HHV}} + \sum (\text{Stack Losses})/m_{AF} \quad (12A)
\]

\[
\text{HPR}_{\text{Act-LHV}} = \text{HPR}_{\text{Ideal-CM-LHV}} + \sum (\text{Stack Losses})/m_{AF} \quad (12B)
\]

\[
\text{HPR}_{\text{Act-HHV}} = \text{HPR}_{\text{Act-LHV}} + \sum n_{\phi} h_{\phi-CAL}^0
\]

\[
+ (n_{\text{H}_2\text{O}} - b_\text{A} - b_z)(\Delta H_{f-CAL-H2O}^0 + h_{\text{Stack}} - h_{\text{g-CAL}})
\]

\[
+ b_\text{A}(1.0 + \beta)(\Delta H_{f-CAL-H2O}^0 + h_{\text{Stack}} - h_{\text{g-CAL}})
\]

\[
+ n_{\text{H}_2\text{O}} z_{\phi}(\Delta H_{f-CAL-CaSO_4}^0 + \Delta H_{f-CAL-H2O}^0 + h_{\text{ESP}} - h_{\text{g-CAL}})
\]

\[
+ b_z[\Delta H_{f-CAL-H2O}^0 + h_{\text{Stack}} - h_{\text{g-CAL}}]/(xN_{AF}) \quad (13)
\]

For non-water products:

\[
HPR_k = [\Delta H_{f-CAL-k}^0 + \int_{T_{\text{CAL}}} dh_k] n_k N_k/ (xN_{AF}) \quad (14)
\]

An absorption efficiency, \(\eta_A\), based on Eq.(11), is derived from HNSL, whose individual terms are evaluated using established procedures:

\[
\text{HNSL} = L_p + L_p + L_{d/Fly} + L_{d/Bott}
\]

\[
+ L_{d/Sorb} + L_{RJ} - W_{fly}/m_{AF} \quad (15)
\]

HNSL bears the same numerical value for both gross or net calculations, as does \(\eta_A\) (see below). All terms of Eq.(15) are specific to unity mass of fuel (i.e., Btu/lbm\(_{AF}\)). Again, these terms affect product streams, no term can be included which does not have such affect. Slight exceptions to standard usage include the following: \(L_p\) is referenced to the total gross (corrected) calorific input, (HHVP + HBC), not HHV or LHV; the \(L_w\) term (of PTC 4.1) is combined with the ash pit term \(L_p\); \(L_{d/Fly}\) is sensible heat in fly ash referenced to its collection point (e.g., the electrostatic precipitator); \(L_{d/Bott}\) is the sensible heat in bottom ash, a separate stream from fly ash; \(L_{d/Sorb}\) is the sensible heat of Sorbent inert matter referenced to its collection point; and \(L_{RJ}\) is the fuel rejects term, an outlet stream.

The ID Fan power, \(W_{ID}\), negatively corrects HNSL such that the fuel’s calorimetrics are conserved. So-called “back-pass” components affect the outlet gas stream by changing boundary temperature and thus the correction (negative for a fan). Most standards advise to leave the ID Fan outside the boundary. This is possible provided temperature and emissions measurements are collected before the fan, which is unusual. It is suggested that all gas fans be included, correcting as indicated. One can argue that correcting a stack temperature for the ID Fan \(\Delta T\) does the same thing. Of course for many situations this might be adequate. However, an exception can be quickly be found with high water systems (e.g., given a high water peat or lignite fuel), in which a 10 \(^\circ\)F correction to stack temperature adds a 0.02% \(\Delta\eta_{B-LHV}\) error given the affect of non-linear water properties.

From Eqs.(11A) and (7A) it becomes apparent that the \(\eta_A\) term is defined as follows:

\[
\eta_A = 1.0 - \frac{\text{HNSL}}{- \text{HPR}_{\text{Act-HHV}} + \text{HRX}_{\text{Act-HHV}}} \quad (16)
\]

There is subtlety here. For consistency with Eq.(11), \(\eta_A\) must be defined in terms of \(\text{HPR}_{\text{Act}}\) and \(\text{HRX}_{\text{Act}}\) and thus \(\eta_A\) becomes a multiplicative affect on boiler efficiency. While this approach minimizes error in \(\eta_B\), the individual terms comprising HNSL have the same weight as stack losses (via \(\text{HPR}_{\text{Act}}\)). In terms of boiler efficiency, Eq.(17) then becomes obvious:

\[
\eta_{B-hHV} = - \frac{- \text{HPR}_{\text{Act-HHV}} + \text{HRX}_{\text{Act-HHV}} - \eta_A}{\text{HHVP} + \text{HBC}} \quad (17A)
\]

\[
\eta_{B-LHV} = - \frac{- \text{HPR}_{\text{Act-LHV}} + \text{HRX}_{\text{Act-LHV}} - \eta_A}{\text{LHVP} + \text{HBC}} \quad (17B)
\]

Note that the “effluent” portion of Eq.(17) is the commonly termed combustion efficiency (i.e., \(\eta_C\), thus, for example: \(\eta_{B-hHV} = \eta_{C-hHV} \eta_A\)).

In summary, all gas/air/working fluid data to the system must be corrected to a common energy level associated with the fuel, \(T_{CAL}\). Firing Corrections affect reactants, while loss terms correct ideal products - both relative to how these terms were established via a
specified $T_{\text{CAL}}$. This is not to say that changes in a Firing Correction might not affect losses; it says that starting from Eq.(4A) lending to Eq.(7), Firing Corrections per se merely balance the calorimetric equation. For the boiler efficiency of Eq.(7), they appear in both numerator and denominator; losses only in the numerator. For ideal combustion, Firing Corrections can not impact efficiency. Without losses, boiler efficiency will always be unity. In the presence of losses, Firing Corrections may impact efficiency but only through the numerics of the efficiency equation. The term HBC is well defined in a previous work (Lang, 2009b).

Air leakage is a fine example of loss effects versus Firing Corrections. If, given an increase in air leakage, Stack temperature is falsely held constant, efficiency will degrade slightly but only because of an increased air flow needed to maintain Boiler $O_2$, but with higher Stack $O_2$ and thus higher losses. However, when lowering Stack temperature reflecting dilution by leakage, one computes a reduction in efficiency.

CALORIFIC CONVERSIONS

Another inconsistency in industrial standards for boiler efficiency lies when converting from gross to net calorific values, and from a constant volume to a constant pressure process. For example, a constant is many times used to convert from constant volume HHV to the constant pressure HHVP (i.e., the As-Fired). Fuel oxygen is sometimes ignored. Underlying correct procedures, the latent heat at constant volume ($\Delta u_{\text{fg,CAL}}$) and the latent heat at constant pressure ($\Delta h_{\text{fg,CAL}}$) at $T_{\text{CAL}}$ must sever as the bases. To convert solid and liquid fuel calorific heating values from a constant volume to a constant pressure process:

\[
\begin{align*}
\text{HHVP} &= \text{HHV} + \Delta h_{\text{V/P}} \\
\text{LHVP} &= \text{LHV} + \Delta h_{\text{V/P}} - \Delta p v_{\text{L/H}} \\
\Delta h_{\text{V/P}} &= RT_{\text{CAL-Abs}} (a_{\text{H}_2}/2 - a_{\text{N}_2} - a_{\text{O}_2}/(JN_{\text{AF}}) \\
\Delta h_{\text{L/H}} &= \Delta h_{\text{fg,CAL-H2O}} (\eta_{\text{ideal-H2O}}) \\
&= \Delta h_{\text{fg,CAL-H2O}} (a_{\text{C}_\text{H}_\text{Gas}}/2 + a_{\text{H}_2} + a_{\text{H}_2S}) N_{\text{H}_2O}/N_{\text{AF}} \\
\Delta u_{\text{L/H}} &= \Delta u_{\text{fg,CAL-H2O}} (a_{\text{C}_\text{H}_\text{Gas}}/2 + a_{\text{H}_2} + a_{\text{H}_2S}) N_{\text{H}_2O}/N_{\text{AF}} \\
\Delta pv_{\text{L/H}} &= \Delta h_{\text{L/H}} - \Delta u_{\text{L/H}}
\end{align*}
\]

where $T_{\text{CAL-Abs}}$ is the absolute. Note that the new ASTM D5828-07 standard employs these same relationships under a work order initially prepared by the author, but employing weight fractions. To convert from a gross (higher) to a net (lower) heating value, the following are exact and consistent with the above:

\[
\begin{align*}
\text{LHV} &= \text{HHV} - \Delta U_{\text{L/H}} \\
\text{LHVP} &= \text{HHVP} - \Delta h_{\text{L/H}}
\end{align*}
\]

The latent heat of water has obvious import when converting from a gross to a net heating value. Although laboratories typically only report the gross, when a fossil fuel is burned its net energy is liberated. There can be no distinction imposed at the time a fuel is burned commercially versus how its energy was determined in the laboratory. If the calorimetric process were to end mid-path, with vaporized water, the derived energy liberated must be the same as would be determined from the completed combustion/vaporization/condensation process. Thus conversion to LHVP serves a useful example. LHVP must employ the $T_{\text{CAL}}$ reference for conversions, to do otherwise implies capricious thermodynamics: one cannot use one latent heat for entrained water, another for water formed from combustion, and yet another latent heat for air’s moisture.

COMPUTED FUEL FLOW, THE ABSOLUTE TEST

Why does adherence to such thermodynamic consistency matter? Is not any definition of efficiency valid if taken as a relative measure (and more so when using high energy fuels)? If an efficiency test produces a boiler efficiency within 0.5 to 1.0% of guarantee, such refinement could matter greatly. If the performance engineer computes an emission flow base on a relative or erroneous boiler efficiency, paying more carbon taxes as a result, it could matter greatly. Such financial arguments aside, industrial standards should be relied on for state-of-the-art methods.

As-Fired fuel flow is not an arbitrary parameter, it is absolute and solely dependent on the fuel’s interaction with gas/air/working fluids (i.e., boiler efficiency). If, indeed, such conversion of fuel energy were dependent (say on an air temperature, an arbitrary 25°C, etc.), then computed fuel flow based on that conversion would be arbitrary. From the Input-Output Method, where $Q_{\text{WF}}$ is the “Useful Energy Flow Developed” of Eq.(1), fuel flow follows by back-calculation:

\[
\begin{align*}
m_{\text{AF}} &= Q_{\text{WF}} / [\eta_{\text{B,HHV}} (\text{HHVP + HBC})] \\
m_{\text{AF}} &= Q_{\text{WF}} / [\eta_{\text{B,HHV}} (\text{HHVP + HBC})]
\end{align*}
\]

When following the guidelines advocated herein, the same fuel flow will be computed given the same losses, using either gross or net efficiency, at any calorimetric temperature and at any level of Firing Corrections. Fuel flow supplied to an in-situ system, interacting with its gas/air/working fluids, is a fixed quantity; it is what it is - it cannot be a function of a user selected reference!!
IDENTITIES

From Eq.(26) it is obvious that the ratio \( \frac{Q_{\text{WF}}}{m_{\text{AF}}} \) is constant, gross and net. Also, from Eq.(11), considering that HNSL and HBC are defined the same for gross and net calculations, the following are developed and considered important identities; see Eq.(16).

\[
\eta_{\text{B-HHV}} (\text{HHVP} + \text{HBC}) = \eta_{\text{B-LHV}} (\text{LHVP} + \text{HBC}) - \eta_{\text{ACT-HHV}} (\text{HHVP} + \text{HBC}) \quad (27)
\]

\[
- \eta_{\text{ACT-HHV}} = - \eta_{\text{ACT-HHV}} + \eta_{\text{ACT-LHV}} \quad (28)
\]

When taking individual stream energies in isolation, the product term becomes irrelevant to gross or net bases as seen in Eq.(13). However, this is also the case for the reactant term given comparison of Eqs.(4B3) and (4C3), and in light of Eq.(25), that:

\[
\text{HHVP} + (\eta_{\text{Ideal-H2O}} \Delta H^0_{\text{g-Cal-H2O}})/(xN_{\text{AF}}) = \text{LHVP} + (\eta_{\text{Ideal-H2O}} \Delta H^0_{\text{g-Cal-H2O}})/(xN_{\text{AF}}) \quad (29)
\]

Thus, as treatment of boundary streams must be divorced from gross or net assumptions, the actual Enthalpy of Product, and Enthalpy of Reactant terms must be identical:

\[
\text{HPR}_{\text{Act-HHV}} = \text{HPR}_{\text{Act-LHV}} \quad (30)
\]

\[
\text{HRX}_{\text{Act-HHV}} = \text{HRX}_{\text{Act-LHV}} \quad (31)
\]

SENSITIVITY TO PROPERTIES

Many would argue that methods of any stripe - be they consistent or tolerated for convenience - are viable if producing “acceptable errors”. However, the problem in agreeing with the notion of an acceptable error is that our responsibility as engineers when writing standards should be to produce state-of-the-art procedures based on sound thermodynamics, one can never anticipate how a standard might be used.

These things said, methods addressing consistent calorimetrics are not needed if testing systems using fuels producing little water on combustion . It is of interest that ASME PTC 4.1 whose pedigree runs back to 1964, with foundations in the Babcock & Wilcox’s Heat Loss Method (taking off another 50 years!), addressed an era in which high energy coal predominated in the U.S.

The source of conceptual calorimetric errors for high product water fuel fuels lies with the sensitivity of the latent heat of water, and its Heat of Formation of the liquid, to temperature as seen in Tables 1A & 1B. The temperature dependency for water’s latent heat is great indicating a 1% sensitivity for every 18 \( \Delta \text{C} \) (10 \( \Delta \text{F} \)). As seen in Table 2 the \( \text{SO}_2 \) sensitivity is slight, while \( \text{CO}_2 \) is 10 times less sensitive than \( \text{SO}_2 \). For fuels with high water contents, the latent heat has obvious import when condensing vapor.

It could be argued that a fossil fuel having low hydrogen but high fuel water has little sensitivity to calorimetric temperature if the bomb’s starting and ending temperature is the same. Under the preceding scenario, could the CV be change? Or, if burning pure graphite in a bomb, if the starting and ending temperature is the same, could the CV be change? The answer is that the CV may very well change due to the path chosen for Eq.(2). By way of example, let’s assume that graphite’s CV is being measured in a calorimeter capable of variable bath temperature. If the bomb is started at 77\(^\circ\)F, then reduced to 34\(^\circ\)F where the graphite is burned, then assume a return to 77\(^\circ\)F - even taking zero change in \( \Delta H^0_{\text{F-CO2}} \) changes reflected in the heat capacity of graphite going to 34\(^\circ\)F, and the heat capacity of \( \text{CO}_2 \) coming from 34\(^\circ\)F, are different resulting in an observed CV which must be different from \( \Delta H^0_{\text{F-CO2}} \) at 77\(^\circ\)F. If water is added to the graphite, path dependency becomes even more obvious. Of course traditional bombs are run with constant temperature in pure \( \text{O}_2 \), and since Eq.(2), and thus Eq.(4A), etc. are path dependent ... we must simply adhere to how the process was run!

The ideal product of combustion, \( \text{HPR}_{\text{Ideal-CF}} \), is defined by the Heats of Formation of the comprising ideal products of: \( \Delta H^0_{\text{F-CO2}} \), \( \Delta H^0_{\text{F-CO2}} \), \( \Delta H^0_{\text{F-CO2}} \), and \( \Delta H^0_{\text{F-CO2}} \). The sensitivity of these terms is reflected in the ratio \( (\Delta \text{HPR}_{\text{Ideal-CF}}/\Delta \text{HPR}_{\text{Ideal-CF}}) \) per 1.0 \( \Delta \text{C} \) as presented in Table 3 for all Ranks of coal. Note that the indicated arrow points to the hvBb Rank fuel (and to those below) whose associated efficiencies can be sufficiently affected by a \( \Delta H^0_{\text{F-CAL}} \) change due to a change in reference temperature. Sensitivity of the \( (\Delta \text{HPR}_{\text{Ideal-CF}}/\Delta \text{HPR}_{\text{Ideal-CF}}) \) term explains why efficiencies associated with the poor quality fuels, having very high product water, have greater sensitivity to \( T_{\text{CAL}} \), and concomitantly have reduced uncertainty versus high energy fuels. Of course this apparent paradox is explained by recognizing the nature of thermochemical properties. Applied thermodynamics of calorimetrics suggests an interesting ambivalence associated with the Energy Balance Method.

When reckoning the insensitivity of \( \text{CO}_2 \), and that seen in Table 3, consider an example of burning pure graphite in a constant temperature bomb. Again let us turn to determining graphite’s CV, not through known Heats of Formation, but rather using a bomb calorimeter. Using results from our graphite calorimeter, one would compute a Energy Balance efficiency which is essentially devoid of temperature dependency. What does it mean then having a system which produces no water and whose Heat of Combustion is insensitive to temperature? Eq.(7) will produce different boiler efficiencies based on any chosen calorimetric bath temperature, using essentially the same CV, but producing a variable fuel flow given “arbitrary” Firing Corrections. Does this imply that our thermo-
dynamic understanding of a graphite system is impossible? No, however it does mean that methods akin to the Energy Balance Method of Eq.(7), are crippled by the nature of thermochemical properties and our inability to discern Heats of Combustion to the accuracy required.

**Simply put, the Energy Balance (Heat Loss) Method has limitations.** Table 3 inherently suggests that a lower limit of 10% by weight of fuel water plus fuel hydrogen be placed on the applicability of the Energy Balance Method.

However, if forced to use an Input-Output approach for low product water fuels, we still have the problem of viable Firing Corrections procedures. For the situation of a fuel having <10% water plus hydrogen, it is proposed to place an error bound on the Input-Output determined boiler efficiency which is dependent on the sensitivity an agreed temperature range has on the Firing Corrections, and thus on boiler efficiency.

Tables 4, 5 and 6 illustrate effects on methane and several coal fuels given a 18 Δ°F (10 Δ°C) change, a misuse, in calorimetric temperature. As seen, differences range from 0.4 to 0.8% in efficiency. The difference between running a bomb at 95°F versus 77°F is not considered uncommon. Nor is it uncommon to compute methane’s CV at 60°F, but then use a 77°F reference. Note that these are only a portion of the conceptual errors. When mis-using loss or “credit” terms, errors have been observed to exceed an additional 1% to 2% in Δη_B (depending on vendor practices and manipulations of standards).

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<td>63.061</td>
<td>+39.99</td>
<td>1102.879</td>
<td>+0.711</td>
<td>1039.818</td>
<td>-0.979</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Reference Temp.</th>
<th>H₂O Heat of Formation,LIQ (Btu/lbm)</th>
<th>PerCent Change (%)</th>
<th>H₂O Heat of Formation,VAP (Btu/lbm)</th>
<th>Change (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>25°C (77°F)</td>
<td>-6821.142</td>
<td>0.000</td>
<td>-5771.140</td>
<td>0.000</td>
</tr>
<tr>
<td>30°C (86°F)</td>
<td>-6817.358</td>
<td>-0.055</td>
<td>-5772.442</td>
<td>+0.023</td>
</tr>
<tr>
<td>35°C (95°F)</td>
<td>-6813.584</td>
<td>-0.110</td>
<td>-5773.766</td>
<td>+0.046</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Reference Temp.</th>
<th>SO₂ Heat of Formation (Btu/lbm)</th>
<th>Change (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>25°C (77°F)</td>
<td>-1992.0031</td>
<td>0.0000</td>
</tr>
<tr>
<td>30°C (86°F)</td>
<td>-1992.1867</td>
<td>0.0092</td>
</tr>
<tr>
<td>35°C (95°F)</td>
<td>-1992.3670</td>
<td>0.0183</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Computed CV for Methane</th>
<th>Gross Efficiency Calculated at 60°F</th>
<th>Gross Efficiency Calculated at 78°F</th>
<th>All Terms / (T_CAL) affecting Δη_B-HHV</th>
</tr>
</thead>
<tbody>
<tr>
<td>23891.00 at 60°F</td>
<td>82.348% (actual)</td>
<td>82.825%</td>
<td>0.477%</td>
</tr>
<tr>
<td>23865.92 at 78°F</td>
<td>82.330%</td>
<td>82.807% (actual)</td>
<td>0.477%</td>
</tr>
</tbody>
</table>
### Table 3: As-Received (wet) Properties of Fossil Fuels
(data from Penn. State, Input/Loss installations & NIST)

<table>
<thead>
<tr>
<th>Substance or Coal Rank</th>
<th>Fuel Hydrogen (%)&lt;sup&gt;wt&lt;/sup&gt;</th>
<th>Fuel Water (%)&lt;sup&gt;wt&lt;/sup&gt;</th>
<th>Combined H&lt;sub&gt;2&lt;/sub&gt; &amp; H&lt;sub&gt;2&lt;/sub&gt;O (%)&lt;sup&gt;wt&lt;/sup&gt;</th>
<th>Fuel Ash (%)&lt;sup&gt;wt&lt;/sup&gt;</th>
<th>Avg. HHV at 25&lt;sup&gt;°&lt;/sup&gt;C (Btu/lbm)</th>
<th>HHV Temp. Coef. (x10&lt;sup&gt;-6&lt;/sup&gt;/1°&lt;sup&gt;°&lt;/sup&gt;C)</th>
<th>ΔHPR&lt;sub&gt;ideal-CF&lt;/sub&gt; HPR&lt;sub&gt;ideal-CF&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>an</td>
<td>1.94</td>
<td>3.55</td>
<td>5.49</td>
<td>9.85</td>
<td>12799.75</td>
<td>19.56</td>
<td>376.6</td>
</tr>
<tr>
<td>sa</td>
<td>3.01</td>
<td>1.44</td>
<td>4.70</td>
<td>16.51</td>
<td>12466.17</td>
<td>30.10</td>
<td>285.0</td>
</tr>
<tr>
<td>lvb</td>
<td>3.97</td>
<td>1.69</td>
<td>5.66</td>
<td>13.22</td>
<td>13155.11</td>
<td>39.22</td>
<td>347.7</td>
</tr>
<tr>
<td>mwb</td>
<td>4.44</td>
<td>1.75</td>
<td>6.19</td>
<td>11.48</td>
<td>13371.75</td>
<td>41.88</td>
<td>380.5</td>
</tr>
<tr>
<td>Benzoic Acid</td>
<td>4.95</td>
<td>0.00</td>
<td>4.95</td>
<td>0.00</td>
<td>11364.57</td>
<td>45.0 (NIST)</td>
<td>---</td>
</tr>
<tr>
<td>hvAb</td>
<td>4.91</td>
<td>2.39</td>
<td>7.30</td>
<td>10.86</td>
<td>13031.61</td>
<td>47.77</td>
<td>444.2</td>
</tr>
<tr>
<td>hvBb</td>
<td>4.63</td>
<td>5.61</td>
<td>10.24</td>
<td>11.83</td>
<td>11852.63</td>
<td>56.53</td>
<td>446.7</td>
</tr>
<tr>
<td>hvCb</td>
<td>4.26</td>
<td>9.89</td>
<td>9.87</td>
<td>12.32</td>
<td>10720.40</td>
<td>60.18</td>
<td>450.6</td>
</tr>
<tr>
<td>subA</td>
<td>3.94</td>
<td>12.93</td>
<td>16.87</td>
<td>7.06</td>
<td>10292.89</td>
<td>51.16</td>
<td>398.3</td>
</tr>
<tr>
<td>subB</td>
<td>3.76</td>
<td>17.87</td>
<td>21.63</td>
<td>9.57</td>
<td>9259.75</td>
<td>61.15</td>
<td>408.0</td>
</tr>
<tr>
<td>subC</td>
<td>3.50</td>
<td>23.79</td>
<td>27.29</td>
<td>10.67</td>
<td>8168.69</td>
<td>75.14</td>
<td>423.3</td>
</tr>
<tr>
<td>ligA</td>
<td>3.02</td>
<td>29.83</td>
<td>32.81</td>
<td>9.64</td>
<td>7294.66</td>
<td>83.56</td>
<td>439.4</td>
</tr>
<tr>
<td>methane</td>
<td>25.33</td>
<td>0.00</td>
<td>25.33</td>
<td>0.00</td>
<td>23867.31</td>
<td>105.39</td>
<td>424.3</td>
</tr>
<tr>
<td>Irish Peat</td>
<td>2.73</td>
<td>46.78</td>
<td>49.51</td>
<td>1.72</td>
<td>4856.07</td>
<td>112.00 (est.)</td>
<td>542.7</td>
</tr>
<tr>
<td>ligB-PSU</td>
<td>2.16</td>
<td>28.84</td>
<td>31.00</td>
<td>22.95</td>
<td>4751.83</td>
<td>122.17</td>
<td>481.3</td>
</tr>
<tr>
<td>ligB-Greek</td>
<td>1.39</td>
<td>54.04</td>
<td>55.43</td>
<td>16.93</td>
<td>2926.82</td>
<td>246.01</td>
<td>685.2</td>
</tr>
</tbody>
</table>

### Table 5: Effects on Gross Boiler Efficiency Given an 18 °F Mis-Use of Calorimetric Temp.

<table>
<thead>
<tr>
<th>Unit</th>
<th>T&lt;sub&gt;CAL&lt;/sub&gt; = 77°F</th>
<th>T&lt;sub&gt;CAL&lt;/sub&gt; = 95°F</th>
<th>Δη&lt;sub&gt;B-HHV&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>110 MWe, CFB, Bituminous Coal Waste</td>
<td>87.678%</td>
<td>88.165%</td>
<td>0.487%</td>
</tr>
<tr>
<td>800 MWe Coal Slurry with Gas Over-Fire</td>
<td>81.174%</td>
<td>81.722%</td>
<td>0.548%</td>
</tr>
</tbody>
</table>

### Table 6: Effects on Gross & Net Boiler Efficiencies Given an 18 °F Mis-Use of Calorimetric Temperature

<table>
<thead>
<tr>
<th>Unit</th>
<th>T&lt;sub&gt;CAL&lt;/sub&gt; = 77°F</th>
<th>T&lt;sub&gt;CAL&lt;/sub&gt; = 95°F</th>
<th>Δη&lt;sub&gt;B-HHV&lt;/sub&gt;</th>
<th>Δη&lt;sub&gt;B-LHV&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>600 MWe PRB Coal, HHV = 8542.828 Btu/lbm</td>
<td>83.602%</td>
<td>84.117%</td>
<td>0.515%</td>
<td></td>
</tr>
<tr>
<td>600 MWe PRB Coal, LHV = 7958.786 Btu/lbm</td>
<td>90.127%</td>
<td>90.646%</td>
<td>0.519%</td>
<td></td>
</tr>
<tr>
<td>300 MWe Lignite-B, HHV = 3369.499 Btu/lbm</td>
<td>68.395%</td>
<td>69.036%</td>
<td>0.641%</td>
<td></td>
</tr>
<tr>
<td>300 MWe Lignite-B, LHV = 2760.249 Btu/lbm</td>
<td>85.047%</td>
<td>85.761%</td>
<td>0.714%</td>
<td></td>
</tr>
</tbody>
</table>
Table 7: Numeric Overchecks Using EX-FOSS

<table>
<thead>
<tr>
<th>Case</th>
<th>Boiler Eff, Eq.(17) %</th>
<th>Fuel Flow, Eq.(26), lb/hr</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HHV</td>
<td>LHV</td>
</tr>
<tr>
<td></td>
<td>HHV</td>
<td>LHV</td>
</tr>
<tr>
<td>Boardman System Evaluation Test, Case SET-Data_Master</td>
<td>84.616</td>
<td>91.688 665,418.39</td>
</tr>
<tr>
<td>Boardman DSI Testing: Trona, Case O2</td>
<td>85.627</td>
<td>92.255 651,387.35</td>
</tr>
<tr>
<td>Boardman DSI Testing: Sodium Bicarbonate, Case G</td>
<td>85.259</td>
<td>91.981 652,061.35</td>
</tr>
<tr>
<td>Graphite Fuel, no losses, T_{CAL} = T_{RA} = T_{Stack} = 77.0°F</td>
<td>100.00</td>
<td>100.00 174,114.02</td>
</tr>
<tr>
<td>Graphite Fuel, no losses, T_{CAL} = T_{RA} = T_{Stack} = 39.0°F</td>
<td>100.00</td>
<td>100.00 174,114.02</td>
</tr>
<tr>
<td>Graphite Fuel, dry air, T_{CAL} = 39.0°F, T_{RA} = T_{Stack} = 77.0°F</td>
<td>98.138</td>
<td>98.138 174,100.33</td>
</tr>
<tr>
<td>PRB Fuel, SET Relative Humidity = 0.0%, T_{Stack} = 278.900°F</td>
<td>84.665</td>
<td>91.742 665,030.08</td>
</tr>
<tr>
<td>PRB Fuel, SET Relative Humidity = 100%, T_{Stack} = 278.900°F</td>
<td>84.611</td>
<td>91.683 665,457.00</td>
</tr>
<tr>
<td>PRB Fuel, SET Relative Humidity = 100%, T_{Stack} = 277.155°F</td>
<td>84.665</td>
<td>91.742 665,030.05</td>
</tr>
</tbody>
</table>

SENSITIVITY TO NUMERICS

Although the author would argue that foundations of Input/Loss Method presented in this work are straightforward (especially with Rev. 30 changes), given the stoichiometric complexity of coal-fired systems, computational over-checks are mandated. An excellent overcheck is to compare Eq.(26A) vs. (26B). Another method of controlling errors is to create a mathematical kernel; for Input/Loss it would consist of Eqs.(4B3), (4C3), (13), (14), (15), HBC calcs, and finally (11), with imbedded properties. Of course basic overchecks should be applied using all standards. Table 7 presents a sampling of EX-FOSS overchecks (Lang, 2012a) for a variety of systems, with and without losses, using different T_{CAL} and reactant conditions; note the matching of fuel flows, HHV versus LHV. For each given case, Firing Corrections (HBC) and \eta_A were computed identically.

Table 7’s last three cases illustrate that although reactant conditions cannot affect ideal combustion, increasing product moisture given a fogged combustion air, while holding T_{Stack} constant, causes higher losses; the process must add heat to bring the air’s moisture to the same T_{Stack}; Stack cooling has been ignored. However, when T_{Stack} is decreased to exactly address such cooling, no change in efficiency or fuel flow is observed.

ASME VERSUS THE WORLD

In North America the gross (higher) heating value based boiler efficiency is the norm for conventional steam generators. In the rest of the world, the net (lower) heating value based efficiency is exclusively used. Net efficiencies are higher numerically. Although for the same fuel, small changes in T_{CAL} have a linear effect on \Delta \eta_B, the affects on \eta_B of changes in product water are quite non-linear. For PRB coals, the \Delta \eta_B impact is approximately the same between gross and net. Note that Table 6 indicates that for very high product water fuels this conclusion breaks down. Although fuel flows will (and must) compute identically for the same T_{CAL} between gross and net, there is numerical subtlety. As \eta_{B-LHV} has the higher value, the percent change of a given \Delta \eta_{B-LHV} is, of course, less versus a percent change in \Delta \eta_{B-HHV}. This condition also implies that the computed fuel flow of Eq.(26B) will be slightly less sensitive to mis-use of T_{CAL}, test uncertainties, etc. The numerical situation implies that greater scrutiny must be given to net-based efficiencies.

SUMMARY AND RECOMMENDATIONS

A consistent boiler efficiency will result if only considering the fuel’s interaction with the gas/air/working fluids, an efficiency leading to a unique fuel flow. Without following such principles, errors can exceed 2% which may have serious effects on power plant monitoring and economics.

The established standards for computing boiler efficiency require improvement. Conceptual errors occur when not adapting the fundamental process of how the energy content of a fossil fuel is determined. In addition,
conceptual errors are made in the treatment of major steam
generator components.

It is the opinion of the author that standards need to
recognize the repeatability accuracy of the modern bomb
calorimeter and, in general, should strive for $\pm 0.1\%$ resolution. The following recommendations, not listed by
importance, are offered for consideration when preparing future revisions of the standards.

1) Boiler efficiency standards should publish, as
integral to their documentation, data sets which can
be replicated by the analyst. Such data sets should
begin with a simple combustion system, with one
heat exchanger, allowing base thermodynamics to
be confirmed and fuel flow computed, followed by
increasing complexity leading to the use of
sorbents, tube leakages, extreme environments, etc.

2) Boiler efficiency standards should be required to
report the reference temperature used in their
computations. This should be the temperature at
which the fuel’s calorific value was determined. If
required to employ a specified reference
temperature, then the coal-fired operator should
specify to the laboratory its desired $T_{\text{CAL}}$.

3) Calorimetric standards such as ASTM D5865 and
ISO 1928:1995(E) should require routine reporting
of calorimetric temperatures. This author, sitting on
the D5865 committee, was defeated in getting
approval of this requirement (his Work Item WK-
5174 was approved by 66 members, with one
negative vote resulting in no modification).

4) Boiler efficiency standards must specify the correct
(and consistent) formula for gross to net, & volume
to pressure conversions, all $T_{\text{CAL}}$ dependent.

5) Standards should specify that any common steam
generator component which was included within
the system boundary (as defined herein), should be
reported and its use fully justified.

6A) Standard committees should investigate the new
NASA combustion gas properties which appear to
have a wider range than the older JANAF tables
(McBride, 2002). For their 2002 publication gas
properties are remarkably consistent with AGA and
CODATA standards. However, there appears to be
some inconsistencies in non-2002 editions such as
NASA’s on-line CEA (Chemical Equilibrium with
Application) data set. All properties should be
normalize to the CODATA standard (Cox, 1989).

6B) It is advised to use water properties based on work
by Keenan, Keyes, Hill and Moore (1969), later
modified by NASA (1973); it represents a truly
consistent and continuous surface fit, resulting in an
excellent matching to international standards.

7) Boiler efficiency standards must anticipate use of
computers and issue approved source coding
(preferably in the simplest of FORTRAN, as was
done for ASME PTC 22).

8) Consistent application should be made of Enthalpy
of Product and Reactant terms as based on the
calorimetric process. At the minimum, the latent
heat and the Heat of Formation for water must be
computed as a function of calorimetric temperature.
All chemistry products should be considered when
enfolding Stack loss terms into a single
formulation. Thus Eq.(13), and a form of Eq.(14),
should be employed thus reducing inconsistencies.

9) For solid and liquid fossil fuels, boiler and
calorimetric standards must not allow corrections
made to as-tested calorific values (other than
conventional conversions from gross to net, etc.).
Such changes to reference temperatures are allowed
under DIN 1942 and are wrong. Laboratory results
based on an as-received sample (HHV) must stand
as-is and be reported, and used uncorrected.

10) Steam generator standards must strictly adhere to
the principle that boiler efficiency can only be
degraded by losses applied to outlet streams. Firing
Corrections are only applied to inlet streams.

11) Modern boiler efficiency standards should allow for
a variable concentration of ambient oxygen. Such
variability was common in older standards, and was
used in test procedures before 1970. Ambient
oxygen is a sensitive parameter for all combustion
equations, fixing this value is ill-advised.

12) The Energy Balance (Heat Loss) Method:
a) should be the only method allowed provided fuel
water plus fuel hydrogen is $\geq10\%$ by weight;
b) evaluations must require the ref. temperature
be equal to the calorimetric temperature;
c) should invoke a tolerance statement on efficiency
based solely on fuel sampling (CV testing) and
those quantities affecting the $Q_{\text{WF}}$ term; and
d) efficiency should be used to back-calculate fuel
flow based on $Q_{\text{WF}}$ (from the Input-Output
definition) for comparison with the actual when-
ever practical and as agreed between parties.
13) The Input-Output Method:
   a) should only be applied if fuel water plus fuel hydrogen is less than 10% by weight;
   b) should use an agreed reference temperature; and
   c) should invoke a tolerance statement on efficiency based on: fuel sampling (CV testing); quantities affecting $Q_{WF}$; and affects on Firing Correction given an agreed spread in reference temperature.

14) It is recommended that standards provide easily understood procedures for uncertainty statements affecting principally fuel sampling and $Q_{WF}$.

15) As ASME PTC 4 is extremely well-detailed, addressing numerous and important testing practices, it is recommended that it serve, after correction of conceptual and procedural errors, as the basis for re-writing European standards.

16) For all ASME standards which determine thermal efficiency, it is recommended that consistent thermodynamic methods and properties be used. Properties would include: molecular weights, air psychrometrics, water properties, gas properties, Heats of Formulation, air’s molar constituents, etc.

NOTES and ERRATA

The following offers historical notes and errata regards this and related works.

Based on numerous comments by colleagues, several minor revisions were made from the original publication in July 2009, through mid-2010; these ending with Rev. 28B.

Starting in November 2010, a formal comparison was made between the methods discussed here and ASME PTC 4 using test data obtained from the 620 MWe coal-fired Boardman unit (operated by Portland General Electric). The initial analyses were reported at the 2011 ASME Power Conference, POWER2011-55215. This testing work resulted in clarifications to this paper, its Revision 29. As reported to the Conference, the Input/Loss analysis did not employ the HPRideal-CM and HRX CAL-CM concepts.

The August 17, 2012 revision (this Rev. 30) represents numerous changes including: 1) a conceptual correction to the base foundation of Input/Loss with changes in nomenclature; 2) adding properties associated with sorbents and their products (Lang, 2012b); 3) use of NASA combustion properties (McBride, 2002); and 4) conversion to Windows 7 with general coding review.

Although the conceptual correction resulting in Eq.(4A) added clarity, changes in PRB-fired efficiencies were typically -0.050% $\Delta \eta_{B-HHV}$ (Rev. 30 being lower). Conceptual corrections also resulted in the identities of Eqs.(30) & (31). Other corrections resulted in $\pm 0.050\%$ to $\pm 0.100\%$ $\Delta \eta_{B-HHV}$.

ACKNOWLEDGMENTS

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Most especially, he would like to thank Mr. Tom Canning (TCanning@TCD.ie), Trinity College, Dublin and formerly with the Electricity Supply Board, Republic of Ireland, for many long hours of discussions and his many critical reviews. The author also gratefully acknowledges his American Society of Mechanical Engineers for allowing his ASME papers to be made available at www.ExergeticSystems.com, they may be downloaded only for personal use.

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Keenan JH, Thermodynamics, Wiley & Sons, New York: 1941; Chapter XVII.


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U.S. Patent 6584429 was issued June 24, 2003. This patent is owned by the author’s company and, upon written request, will be made available for ASME’s use without royalties; use outside the U.S. requires no request.


Vakkilainen EK and Ahtila P, “Modern Method to Determine Recovery Boiler Efficiency”, 2009 TAPPI Engg., Pulping, Environmental Conference”, from: www.tappi.org/Bookstore/Technical-Papers/Conference-Papers/2009/09EPE.aspx. This paper states “Recovery boiler mass and energy balances are needed for performance testing of recovery boilers … [however] the use of [TAPPI,1996] is problematic in Europe because of its view that losses from recovery of process chemicals are counted when determining the recovery boiler steam generation efficiency. Low efficiency leads to taxation and legislative problems in some European countries. … This paper examines how [EN 12952-15:2003] can be applied to recovery boilers to find out net efficiency which is to be approximately [the] same [as] for other biomass boilers, not 10 % lower.”

“Water-Tube Boilers and Auxiliary Installations - Part 15: Acceptance Tests”, prEN 12952-15: November 1999, European Committee for Standardization, Brussels (also: CEN/TC 269/WG 3N 337); the draft European standard (reviewed in 2002); now EN 12952-15:2003 replacing the German and British standards.