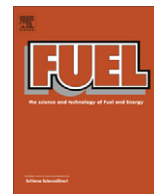




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A laminar burning velocity and flame thickness correlation for ethanol–air mixtures valid at spark-ignition engine conditions

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HIGHLIGHTS

- We evaluate laminar burning velocity and flame thickness correlations for ethanol.
- Reliable ethanol laminar burning velocity data at engine conditions are scarce.
- We calculate laminar ethanol flames at engine conditions using chemical kinetics.
- A new laminar burning velocity correlation form closely fits the calculation results.
- An existing flame thickness correlation is adapted to be valid for ethanol flames.

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ABSTRACT

The use of biomass-derived ethanol in spark-ignition engines is an interesting option to decarbonize transport and increase energy security. An engine cycle code valid for this fuel, could help to explore its full potential. Crucial building blocks to model the combustion in ethanol engines are the laminar burning velocity and flame thickness of the ethanol–air–residuals mixture at instantaneous cylinder pressure and temperature. This information is often implemented in engine codes using correlations. A literature survey showed that the few available flame thickness correlations have not yet been validated for ethanol. Also, none of the existing ethanol laminar burning velocity correlations covers the entire temperature, pressure and mixture composition range as encountered in spark-ignition engines. Moreover, most of these correlations are based on measurements that are compromised by the effects of flame stretch and the occurrence of flame instabilities. For this reason, we started working on new correlations based on flame simulations using a one-dimensional chemical kinetics code.

In this paper the published experimental data for the laminar burning velocity of ethanol are reviewed. Next, the performance of several reaction mechanisms for the oxidation kinetics of ethanol–air mixtures is compared. The best performing mechanisms are used to calculate the laminar burning velocity and flame thickness of these mixtures in a wide range of temperatures, pressures and compositions. Finally, based on these calculations, correlations for the laminar burning velocity and flame thickness covering the entire operating range of ethanol-fueled spark-ignition engines, are presented. These correlations can now be implemented in an engine code.

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1. Introduction

Rising fuel prices, air pollution and the consequences of global warming make the need for a sustainable alternative for fossil fuels painfully clear. The use of biomass-derived ethanol in spark-ignition engines is receiving increasing attention these days, since it is an interesting option to decarbonize transport and increase energy security. Compared to other promising alternatives, such as hydrogen and battery electric vehicles, liquid ethanol entails

less issues regarding transport and distribution and is easily stored in a vehicle. In addition, it can be blended with standard gasoline in variable amounts, facilitating a gradual transition to renewable fuels.

Bio-ethanol blends can be used in low-cost spark-ignition engines with only minor adjustments. Unlike many other alternative fuels, they have the potential to increase the engine performance and efficiency while reducing emissions compared to gasoline. This is especially true for blends with high ethanol content (e.g. E85, E100). An interesting way to expedite evolution towards these high concentration blends is to use them in flexible fuel vehicles. These engines give the driver the choice what fuel to use, but their

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performance is usually a compromise between operation on gasoline or alcohol.

State-of-the-art flex-fuel vehicles utilize the evaporative cooling effect of alcohols to suppress knock in highly charged engines with direct injection. Saab reports that such engines can attain 20% more power on E85, whereas the mean brake thermal efficiency over the New European Driving Cycle improves by over 5% compared to operation on gasoline [1].

Alternatively, the favourable characteristics of ethanol can be fully exploited in dedicated engines. These engines produce high levels of power and efficiency through the use of design measures such as high compression ratios and downsizing, but this makes them unfit for operation on gasoline. To illustrate their potential, researchers at the U.S. Environmental Protection Agency converted a production 1.9 litre turbocharged diesel engine with a compression ratio of 19.5:1 to run on neat ethanol in SI mode and were able to attain peak effective thermal efficiencies higher than the baseline diesel engine (>40%) [2]. EGR levels up to 50%, made possible through the high dilution tolerance of ethanol, were used to spread the high efficiency regions to part-load operating points. These experimental results illustrate the potential of ethanol. A more comprehensive discussion on the favourable properties of alcohols and their use in spark-ignition engines can be found in [3,4].

Despite these promising experimental results, the full potential of bio-ethanol in modern engine technology and its impact on possible control strategies remain to be explored. With current trends like elevated pressure-charging (downsizing), variable valve timing, exhaust gas recirculation, etc. it is no longer possible for an R&D engineer to intuitively grasp how these technologies and their combinations will affect the engine performance and pollutant emissions. Today however, these issues can be addressed at low cost using system simulations of the whole engines, provided that the employed models account for the effect of the fuel on the combustion process.

Predictive engine codes require a turbulent combustion model to track the progress of the flame front through the cylinder. These combustion models generally assume fast chemistry (i.e. flamelet combustion regime). The influence of turbulence is then limited to flame stretching and the increase of flame front area, whereas the contribution of chemical reactions is grouped in the laminar burning velocity. This is a physico-chemical property of the fuel–air–residuals mixture and thus a fundamental building block of any engine model. Another important flame property is the laminar flame thickness δ_f . Several recent turbulent combustion models use this property to estimate the ability of different turbulence scales to wrinkle the flame front and strain the flame [5–7]. This is done to reflect the observation that compared to thick flames, thin flames can be wrinkled by smaller turbulent length scales, thus producing more surface area and higher turbulent burning velocities.

Both laminar burning velocity and flame thickness are dependent on pressure, unburned mixture temperature and composition. Therefore turbulent combustion models require their values at instantaneous in-cylinder conditions. A convenient way to implement laminar burning velocity and flame thickness data in an engine cycle code is by using a correlation that gives their values in terms of pressure, temperature and composition of the unburned mixture. Unfortunately, none of the published laminar burning velocity correlations covers the entire range of conditions as encountered in ethanol-fueled spark-ignition engines. Their validity is also doubtful, since most of them are based on outdated measurements which do not account for the effects of flame stretch and instabilities on the burning velocity. For flame thickness δ_f different fuel-independent correlations have been proposed, but these have not yet been validated for ethanol–air flames.

The present work seeks to evaluate the existing correlations for the laminar burning velocity and flame thickness of ethanol–air

flames. It was completed within the framework of developing a multi-zone thermodynamic engine code valid for alcohol-fueled engines. A first part of the paper reviews the published data on ethanol laminar burning velocity. We evaluated to what extent the current correlations cover the operating range of ethanol-fueled engines and selected the most reliable experimental results. Besides reviewing experimental work, we also examined different reaction mechanisms for the oxidation kinetics of ethanol–air mixtures. A series of calculations were performed using the one dimensional chemical kinetics code CHEM1D [8] in order to determine which mechanisms could best represent the experimental values for the laminar burning velocity. Using these selected mechanisms, we calculated the burning velocity of ethanol–air mixtures under a broad range of mixture compositions, pressures and temperatures. Finally, based on this set of calculations, the performance of existing laminar burning velocity and flame thickness correlations is evaluated and improved correlations are proposed.

2. Choice of chemical kinetic scheme

2.1. Experimental determination and numerical calculation of laminar burning velocity

Most published laminar burning velocity correlations are based on a limited set of experimental data. Table 1 summarizes the major experimental investigations of ethanol laminar burning velocity. An in-depth review of the employed measurement methodologies and reliability of the results was published in [9]. The main conclusions are repeated here and some more recent experimental data and insights are added.

In [9] the in-cylinder conditions of future flex-fuel and dedicated ethanol engines are defined as in Table 2. When comparing this to the experimental domains in Table 1, it can be seen that the validation range of current correlations [10,11] is rather limited. Data at elevated pressure is scarce and none of the investigations look at high levels of dilution.

Moreover, most of the older u_f data is compromised by the effects of flame stretch and instabilities. For example, Gülder [10], who published one of the earliest extensive measurement sets, investigated the flame propagation of contained explosions in a closed vessel. This is a popular way to measure u_f at elevated pressures. The flame propagation can either be derived from pressure measurements in the vessel [10,12] or directly captured by a high speed camera and a Schlieren optical system [11,13–16]. The stretch due to the curvature of these spherical flames can lead to substantial under or overestimations of the real burning velocities if not corrected for [10,15]. Additionally the spherical flames can develop instabilities such as cellularity at high pressures and temperatures. These can cause overestimations of the true burning velocity at these conditions [10]. Gülder ignored these two effects in his research. Consequently his correlation can be expected to give incorrect u_f values. In more recent publications, the effects of stretch and instabilities are accounted for. Still, the right method to correct for flame stretch remains subject to debate [17,18].

As mentioned above, experimental determination of laminar burning velocities at engine-like conditions is hampered by the occurrence of flame instabilities and incorrect stretch corrections cause scatter amongst published data. Computationally, these effects can be avoided by assuming one-dimensional, planar adiabatic flames. The accuracy of burning velocities calculated with this assumption then depends on the correctness of the chemical kinetic reaction scheme and the precision of the rate constants and molecular transport coefficients. Understandably, the validation of reaction mechanisms against laminar burning velocities is very limited at best. Most mechanisms are therefore more widely

Table 1

Overview of the ethanol–air burning velocity measurements in literature. * Correlation published. There is a lack of data at elevated pressures and for diluted mixtures.

Year	Author	Ref.	Technique	p (bar)	T (K)	ϕ	f (m%)
1982	Gülder*	[10]	Closed vessel, flame ionization	1–8	298–800	0.7–1.4	0
1992	Egolfopoulos et al.	[26]	Counterflow	1	363–453	0.55–1.8	0
2006	Liao et al.*	[11]	Closed vessel Schlieren	1	385–480	0.8–1.2	0
2006	Kimitoshi et al.	[13]	Closed vessel Schlieren	1	325	0.8–1.4	0
2009	Bradley et al.	[14]	Closed vessel Schlieren	1–14	300–393	0.7–1.5	0
2009	Ohara et al.	[15]	Closed vessel Schlieren	1–5	298	0.8–1.4	0
2009	Beeckmann et al.	[16]	Closed vessel Schlieren	10	373	0.8–1.2	0
2010	Veloo et al.	[18]	Counterflow	1	343	0.7–1.5	0
2010	Konnov et al.	[25]	Flat flame heat-flux method	1	298–358	0.65–1.55	0
2011	Eisazadeh-Far et al.* (Far)	[12]	Closed vessel pressure derived	1–5	300–650	0.8–1.1	0–10
2011	Broustail et al.	[35]	Closed vessel Schlieren	1	393	0.8–1.4	0
2011	Varea et al.	[29]	Closed vessel Schlieren	1–5	373	0.8–1.5	0
2011	Marshall et al.	[28]	Closed vessel pressure derived	0.5–4	400–650	0.7–1.4	0–30

Table 2

In-cylinder conditions in alcohol-fueled engines.

Engine type	p (bar)	T (K)	ϕ	EGR (m%)
Flex-fuel	1–100	300–1000	0–2.5	0–20
Dedicated	1–150	300–1500	0–2.5	0–50

validated. Typically they are tested on the basis of measured ignition delay times, flame extinction stretch rates, concentration profiles from flow reactors and flame data from burners. The accuracy of such a comprehensive mechanism is then a trade-off between the several applications it was developed for.

Several researchers have developed comprehensive mechanisms for the oxidation kinetics of ethanol–air mixtures (see Table 3). In order to determine which mechanisms are most fit to calculate laminar burning velocities under engine-like conditions, a number of simulations were run with a one-dimensional chemical kinetics code (CHEM1D) [8]. This code was developed at Eindhoven University of Technology and employs the EGLIB complex transport model [19], including multicomponent transport and thermal diffusion. In each case the solution was calculated using the exponential differencing technique in a grid consisting of 200 points, with most of the detail centred at the inner flame layer. Radiation was neglected and solver convergence confirmed by ensuring that all residuals were below 10^{-10} and the laminar burning velocity had reached a stable value. A grid independence test was performed to eliminate the large truncation errors from inadequate grid resolution. It was shown that the laminar burning velocity differed by less than 1% between 200 and 400 grid points.

2.2. Comparison of experimental and numerical results

2.2.1. Laminar burning velocity at varying equivalence ratio

The predictive performance of various ethanol oxidation mechanisms is tested for premixed flames at 358 K, 1 bar and for varying equivalence ratio in Fig. 1. The mechanisms of Marinov [20], Li et al. [21], Saxena and Williams [22] and Konnov et al. [23] are included in the comparison. These are some of the more recent and widely validated ethanol mechanisms available [9]. Röhl and

Peters [24] have published a reduced version of the Marinov mechanism that gives almost identical results to the original scheme (not shown here).

The experimental datasets of Bradley et al. [14], Konnov et al. [25] and Egolfopoulos et al. [26] are chosen as point of reference. Bradley et al. [14] performed the most extensive contemporary experimental investigation of ethanol–air burning velocity. They captured the flame growth of contained explosions using a high speed camera and a Schlieren optical system. A linear extrapolation to zero stretch was performed after removing all cellular flames and flames over-driven by spark energy from the dataset. Konnov et al. [25] used an interesting alternative set-up to measure the laminar burning velocity of ethanol–air mixtures at atmospheric pressure. They used a flat flame burner to stabilize non-stretched flames on a perforated plate burner. The so-called heat flux method was used to determine the burning velocity under conditions where the net heat loss from the flame to the burner is zero. The overall accuracy on the measured burning velocities was estimated to be better than 1 cm/s. Note the strong correspondence between these two recent datasets, although the authors gathered their data using completely different measurement methods. Van Lipzig et al. [27] partly repeated the measurement set of Konnov et al. [25] using an identical setup. Their experimental values were systematically higher, which was traced back to an incorrect placing of the outer thermocouple on the perforated plate burner in the work of Konnov et al. The difference in maximum burning velocity was about 2 cm/s and for lean flames the divergence mounted to 3–4 cm/s. Egolfopoulos et al. [26] employed a counterflow twin-flame burner to measure the burning velocity of ethanol for a range of equivalence ratios (0.5–2) at atmospheric pressure and modest temperatures (363–453 K). Because the typical strain rate in their flames is quite small (about 100 s^{-1}), they used a linear extrapolation to zero stretch, which was later reported to lead to overestimations of the true burning velocity of up to 10% [18]. On Fig. 1 it can be seen that the u_l values of Egolfopoulos et al. are consistently higher than the other datasets, especially around stoichiometry.

Also plotted in Fig. 1 are the expressions of Liao et al. [11] and Eisazadeh-Far et al. [12] to demonstrate the predictive capabilities

Table 3

Overview of the employed ethanol oxidation mechanisms and their validation range.

Year	Author	Ref.	Species	Reactions	Validation	p (bar)	T (K)	ϕ
1998	Marinov (Mar.)	[20]	57	383	u_l shock tube flow reactor	1–4.5	300–1700	0.5–2.0
2006	Li et al.	[21]	39	238	u_l shock tube flow reactor	1–13	300–950	0.3–1.4
2007	Saxena and Williams (S&W)	[22]	57	288	u_l shock tube flame struct.	<100	>1000	<3
2009	Röhl and Peters	[24]	38	228	u_l flow reactor	1–40	300–800	0.5–2
2009	Konnov	[23]	127	1200	u_l	1	298–358	0.5–2

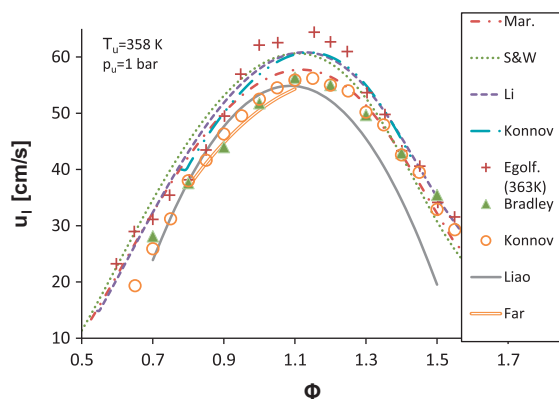


Fig. 1. u_l of ethanol–air as a function of ϕ ($p = 1$ bar, $T_u = 358$ K). Comparison of chemical kinetic mechanisms (discontinuous lines), experimental data (markers) and existing correlations (full lines). References in Tables 1 and 3.

of current laminar burning velocity correlations. Liao et al. [11] founded their correlation on measurements gathered using a similar method as Bradley et al. Eisazadeh-Far et al. also employed contained explosions, but derived the burning velocity using pressure measurements and a multi-zone burning model. Only stable flames with radii higher than 4 cm (corresponding to stretch rates less than 70 s^{-1}) were retained in their dataset to exclude the effects of flame stretch and instabilities. This restricted the usable data to equivalence ratios between 0.8 and 1.1, temperatures between 300 and 650 K and pressures below 5 bar. Both correlations predict burning velocities comparable to the values of Bradley et al. and Konnov et al. for equivalence ratios below 1.1. For rich mixtures the correlation of Liao et al. gives slightly lower values.

When comparing the calculation results using the chemical kinetic mechanisms to the experimental burning velocity values, it can be seen that for lean to stoichiometric mixtures all schemes predict higher burning velocities than the values reported by Konnov et al. and Bradley et al. Keeping in mind the observations of Van Lipzig et al. [27] one can expect the results of Konnov et al. to be 2–4 cm/s too low. For slightly rich mixtures the mechanism of Marinov gives the best match with the data of Konnov et al. and Bradley et al. For very rich mixtures there is not much to choose between the four mechanisms under consideration.

2.2.2. Laminar burning velocity at varying unburned mixture temperature

Fig. 2 shows the laminar burning velocities for a range of unburned mixture temperatures and for stoichiometric mixtures at atmospheric pressure. Compared to Fig. 1 the individual data points of Liao et al. [11] and the correlation of Marshall et al. [28] are added. Similar to Eisazadeh-Far et al., Marshall et al. derived the burning velocity using pressure measurements and a multi-zone burning model. Cellular flames were removed from the dataset after visual inspection of Schlieren photographs of the flame front. The readings will thus be influenced by the criteria used to define the onset of cellularity. No stretch correction was applied, but the authors claim that the mean error due to stretch should be around 2 cm/s. However, it is difficult to estimate to what extent measurements at low and high temperatures are equally affected. From Fig. 2 it can be seen that for low to intermediate temperatures the difference between the mechanisms remains negligible. For temperatures exceeding 500 K the mechanism of Marinov predicts significantly lower values (10%) than the other mechanisms. Looking at the correlations of Liao et al. and Eisazadeh-Far et al. the temperature evolution predicted by the Marinov scheme seems more trustworthy. However, the data points gathered by Liao et al. and

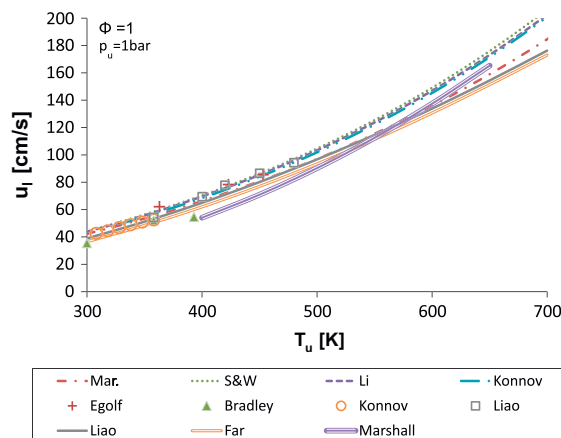


Fig. 2. u_l of ethanol–air as a function of T_u ($p = 1$ bar, $\phi = 1$). References in Tables 1 and 3.

the correlation of Marshall et al. support the steeper temperature gradient. Actually, the temperature range of the available measurement data is too restricted and the uncertainty on measurements too large to draw any sound conclusion on the temperature behavior of the various mechanisms.

2.2.3. Laminar burning velocity at varying pressure

The dataset of Ohara et al. [15] and especially that of Bradley et al. [14] provide interesting information on the pressure behavior of ethanol–air flames. Fig. 3 compares the calculated u_l against these datasets for varying pressure at 353 K and for three different equivalence ratios. Also included are the correlations of Liao et al. [11] and Eisazadeh et al. [12], and the dataset of Varea et al. [29] at $\phi = 1$ and $T_u = 373$ K. Similarly to Bradley et al., Varea et al. obtained the burning velocity by capturing the flame growth during contained explosions.

The dataset of Ohara et al., Bradley et al. and Varea et al. suggest an exponential decrease in burning velocity with increasing pressure. For this reason the results from the chemical kinetic calculations are displayed on a logarithmic coordinate system. It is clear that the current correlations underpredict the drop in burning velocity with increasing pressure. The pressure dependence of the correlation of Liao et al. is based on measurements of Gülder et al. [10], which possibly overestimate the true burning velocity of high pressure flames due to the inclusion of cellular flames. Eisazadeh et al. excluded cellular flames from their dataset, but this left them with a dataset that was very limited in pressure range (<4 bar at $\phi = 1$).

For lean to stoichiometric mixtures, all mechanisms predict a similar pressure decrease as measured by Bradley et al., Ohara et al. and Varea et al. Note that the results of Varea et al. are measured at 373 K and therefore somewhat higher. As mentioned above, the calculated u_l values are somewhat higher. For rich mixtures the scheme of Marinov predicts a steeper decrease in u_l than the other schemes. One could see a match between the calculations of Marinov and the measurements of Ohara et al., but the data of Ohara et al. was gathered at a lower temperature and cannot be directly compared. The experimental values of Bradley et al. support the calculations of Li et al. and Saxena and Williams. Although not shown here, the mechanism of Marinov and its reduced equivalent also underestimate the laminar burning velocity at elevated pressures for other equivalence ratios on the rich side.

2.2.4. Laminar burning velocity at varying diluent ratio

To the author's knowledge the only experimental works that looked at the burning velocity of diluted ethanol–air flames were

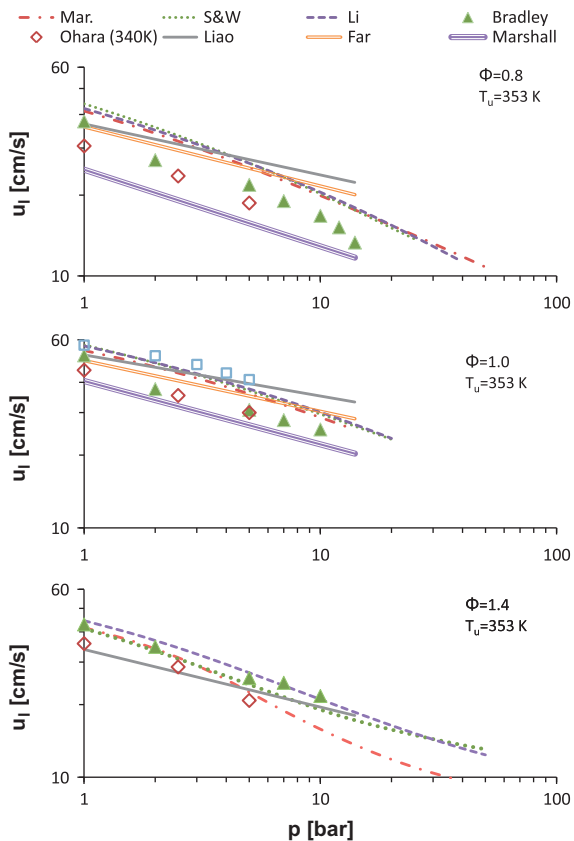


Fig. 3. u_l of ethanol–air as a function of p at different equivalence ratios ($T_u = 353$ K). References in Tables 1 and 3.

published by Eisazadeh et al. [12] and Marshall et al. [28]. The former used a mixture of 86% nitrogen and 14% carbon dioxide to simulate the heat capacity of residual gases in concentrations up to 10% by volume. The latter employed a preliminary explosion and left part of the residuals in the bomb to investigate the effect of residuals in concentrations up to 30% by volume. Gülder's correlation also has a term to include the effect of diluents, but it was based on methanol–air experiments and not validated for ethanol–air mixtures. Fig. 4 compares the correlation of Eisazadeh et al. and Gülder with the values predicted by different chemical kinetic schemes. As can be seen from the figure, the different mechanisms predict a similar decrease in burning velocity with

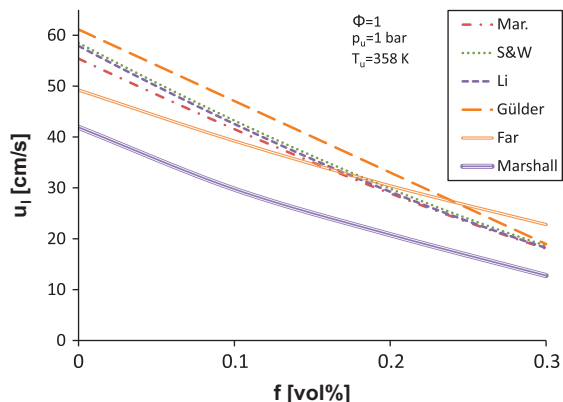


Fig. 4. u_l of ethanol–air as a function of f ($p = 1$ bar, $T_u = 358$ K, $\phi = 1$). References in Tables 1 and 3.

increasing diluent ratio. Surprisingly, the linear decrease predicted by the correlation of Gülder is quite close to the calculation results. The correlation of Eisazadeh et al. and Marshall et al. seem to indicate that the influence of dilution on u_l is less pronounced. However, it must be noticed that their correlations were based on measurements at higher temperatures, which have been shown to be less affected by residuals [30].

Based on the results discussed in the previous paragraphs, the mechanism of Li et al. [21] was chosen to calculate the burning velocity of ethanol–air mixtures under a wide range of temperatures (400–1000 K), pressures (5–105 bar), equivalence ratios ($0.5 \leq \phi \leq 2$) and diluent ratios (0–50 vol.%) as they appear in spark-ignition engines. It is a contemporary, widely validated mechanism that is computationally not too demanding and corresponds well with the published measurements under the conditions examined. The reduced version of the Marinov scheme [20] published by Röhl and Peters [24] is an interesting alternative, especially when one is not interested in rich mixtures at elevated pressures. More experimental laminar burning velocity data at elevated pressures and dilution ratios remain desirable, however, to further validate these reaction schemes.

3. Laminar burning velocity correlation

3.1. Evaluation of published correlations

Based on the calculated flame database of over 1500 conditions, the existing laminar burning velocity and flame thickness correlations of ethanol–air mixtures can be evaluated. As shown in Table 1 only a few laminar burning velocity correlations for ethanol–air mixtures were published. All of these use the form shown in Eq. (1) to express the influence of equivalence ratio, pressure, unburned mixture temperature and residual gas content on the burning velocity. This form has frequently been used for various fuels.

$$u_l = u_{l0} \cdot (T_u/T_{u0})^\alpha \cdot (p/p_0)^\beta \cdot (1 - \gamma f) \quad (1)$$

where p_0 , T_{u0} and u_{l0} are the pressure, unburned gas temperature and laminar burning velocity at atmospheric reference conditions. A second-order polynomial in ϕ is often used to fit u_{l0} . The pressure and temperature exponents α and β are given by linear functions of ϕ . The amount of residuals in mass fraction is represented by f . Its coefficient γ is usually constant.

Analysis of our calculation data for ethanol–air flames supported the exponential trends in pressure and unburned mixture temperature. However, whereas Eq. (1) assumes the effects of ϕ , p , T_u and f to be independent, our analysis revealed there can be a strong interaction between the effects of ϕ , p and T_u . To illustrate this, Fig. 5 shows the calculated temperature exponent α of ethanol–air flames for a range of pressures and equivalence ratios. Note how α rises for increasing pressure at lean equivalence ratios and decreases as a function of pressure at rich equivalence ratios. From this Figure it can also be seen that the reduction of α with increasing equivalence ratio as reported in some works [11,12] is not reflected in the results of our calculations. Similar interactions can be seen when the pressure exponent β is plotted as a function of equivalence ratio and unburned mixture temperature (not shown here). The results from our calculation indicate that β is an increasing function of ϕ , even for rich mixtures. Eisazadeh et al. propose a linearly increasing function of ϕ . In contrast, Gülder's pressure exponent β is an increasing function of ϕ for lean mixtures, but decreases linearly with equivalence ratio for rich mixtures.

Verhelst et al. [30] noted the same strong interaction effects in their calculations results for the laminar burning velocity of hydrogen–air mixtures. To cover these effects, they propose the following functional form for the correlation:

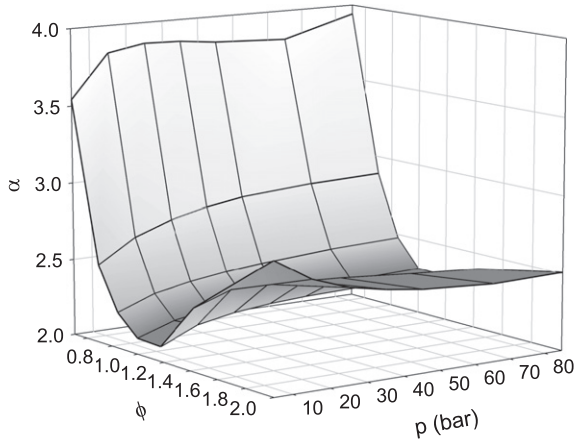


Fig. 5. Calculated temperature exponent α of ethanol–air mixtures for a range of pressures and equivalence ratios. The calculated pressure behavior of lean flames and rich flames is different.

$$u_l(\phi, p, T_u, f) = u_{l0}(\phi, p) \left(\frac{T_u}{T_0} \right)^{\alpha(\phi, p)} F(\phi, p, T_u, f) \quad (2)$$

where both $\alpha(\phi, p)$ and $u_{l0}(\phi, p)$ are polynomial functions of ϕ and p with cross-terms due to the strong interaction between these variables. Verhelst et al. also suggest that the correction term F to account for residual gases is a complicated function of ϕ , p , T_u and f . From the calculation results presented in Fig. 4 it could already be seen that the linear decrease in u_l with increasing diluents is only valid for low diluents fractions (<10 vol.%). Inspection of the calculated correction factor F in function of ϕ , p , T_u and f shows that the dominating factor is the diluent volume fraction, but there are indeed important effects of T_u , p and ϕ . For example, the tolerance for dilution rises at elevated temperatures. This effect of T_u on the correction factor F is more pronounced for rich mixtures and at elevated pressures. Similar interaction effects were reported by Verhelst et al. for the laminar burning velocity of hydrogen–air mixtures [30].

3.2. New laminar burning velocity correlation

For reasons mentioned above, the functional form proposed by Verhelst et al. was selected for the correlation (Eq. (2)). To make the pressure and temperature non-dimensional the standard reference conditions were used ($p_0 = 1$ bar, $T_0 = 300$ K).

For undiluted mixtures the exponent α and coefficient u_{l0} of the power relation were determined at each combination of ϕ and p . However, inspection of the dataset for ethanol–air mixtures revealed that the proposed power relation only holds for temperatures below 900 K. For temperatures above 900 K and at elevated pressures, there were substantial deviations. This behavior is due to the fact that at these temperatures the mixture self-ignites and the definition of laminar burning velocity has no physical sense (the self-ignition temperature of ethanol is 698 K at atmospheric conditions). Consequently, the calculation results at the highest pressures (>85 bar) and temperatures (≥ 900 K) were removed from our dataset.

The Levenberg–Marquardt algorithm [31] was used to fit the calculated α and u_{l0} values as a function of ϕ and p . This algorithm seeks to reduce the sum of the squared differences (SSD) between the observed and predicted values. Due to the large spread in u_l values (ranging from a few cm/s to more than 2 m/s) a weighting parameter was used during the fitting, to ensure an accurate fit also for the lower burning velocities. The weighting was set to the squared reciprocal of the observed value as this gave the best

results. Each of the proposed equations (see below, Eqs. (3) and (4)) in this paper is the result of a large number of iterations, whereby different functional forms were fitted to the data. Initially these forms consisted of only linear terms in the different variables ($p/p_0, \phi, f, T/T_0$). Progressively terms were added, first pure quadratic terms ($(p/p_0)^2, \phi^2, f^2, (T/T_0)^2$) followed by linear cross-terms (e.g. $(p/p_0) \cdot (T/T_0), (T/T_0) \cdot f, \dots$), inverse linear terms (e.g. $1/(p/p_0)$) and progressively higher order terms and combinations of these factors. This was continued until the resulting SSD no longer decreased. Once this stage was reached it was attempted to trim the equation, by selectively removing terms one by one to see their impact on the SSD. The resulting fit was always visually compared to the original simulation data to confirm the effect of each parameter was well represented by the correlation form.

Table 4 lists the coefficients to determine $\alpha(\phi, p)$ using Eq. (3) and $\ln[u_{l0}(\phi, p)]$ using Eq. (4). Due to the large variation in u_{l0} values it was decided to fit $\ln u_{l0}$ in order to reduce the complexity of the fit. Note that Eq. (4) is third order in p , whereas the burning velocity decreases exponentially with increasing pressure. This leads to underestimated burning velocities at the lowest pressures (<10 bar). These points are not that important considering the intended purpose of the correlations in an engine simulation code.

$$\alpha(\phi, p) = a_1 + a_2\phi + a_3\frac{p}{p_0} + a_4\phi\frac{p}{p_0} + a_5\phi^2 + a_6\left(\frac{p}{p_0}\right)^2 + a_7\phi^3 + a_8\left(\frac{p}{p_0}\right)^3 + a_9\phi^2\frac{p}{p_0} + a_{10}\phi^2\left(\frac{p}{p_0}\right)^2 + \frac{a_{11}}{\phi} + \frac{a_{12}}{\phi}\frac{p}{p_0} \quad (3)$$

$$\ln[u_{l0}](\phi, p) = b_1 + b_2\phi + b_3\frac{p}{p_0} + b_4\phi\frac{p}{p_0} + b_5\phi^2 + b_6\left(\frac{p}{p_0}\right)^2 + b_7\phi^3 + b_8\left(\frac{p}{p_0}\right)^3 + b_9\phi^2\frac{p}{p_0} + b_{10}\phi^2\left(\frac{p}{p_0}\right)^2 + \frac{b_{11}}{\phi} \quad (4)$$

Once the correlation for undiluted mixtures is known, an expression for residual gas correction term can be constructed. As mentioned above, the best way to capture the influences and interactions of ϕ , p , T_u and f on the correction term F , is by fitting this term as a complicated polynomial function. The data series for the correction term is computed as the ratio of the calculated dataset values, with residuals, to the corresponding values without residuals, predicted using the correlation proposed above (Eqs. (2)–(4), $F = 1$). Using a similar procedure as described above a functional form was determined for the correction term, minimizing the SSD. During this fitting procedure the smallest u_l values (<1 cm/s) were removed from the dataset. The lowest burning velocities occur for extreme equivalence ratios (0.5, 2.0) at the highest concentrations of residual gases (>30 vol.%). These flames might exist computationally, but are not considered appropriate for engine

Table 4
Coefficients for Eqs. (3) and (4).

i	a_i	b_i
1	3.717600E+00	−3.123300E+00
2	−9.398400E+00	2.054070E+01
3	3.980000E−02	−5.880000E−02
4	−1.860000E−02	9.329600E−03
5	8.413800E+00	−1.617230E+01
6	−2.832200E−04	7.285600E−04
7	−2.055000E+00	3.633400E+00
8	1.401100E−06	−3.097700E−06
9	−8.349800E−04	4.372200E−03
10	4.319800E−05	−1.179500E−04
11	1.332500E+00	−1.434600E+00
12	−6.523800E−03	−

Table 5
Coefficients for Eq. (5).

i	c_i	i	c_i
1	1.087600E+00	11	5.054000E-01
2	1.088600E+00	12	1.851200E+00
3	-4.133000E-01	13	1.803000E-01
4	-2.787700E-03	14	-2.108700E+00
5	-6.703000E+00	15	1.680000E-02
6	-7.413000E-01	16	-5.028100E-04
7	1.250000E-01	17	4.040000E-02
8	8.049200E+00	18	-2.813000E-01
9	-2.486000E-01	19	-7.175000E-01
10	3.037800E-03	20	-1.323600E+00

simulations. Table 5 lists the coefficients to determine $F_1(\phi, p, T_u, f)$ using Eq. (5). The correction term $F(\phi, p, T_u, f)$ is then found by limiting $F_1(\phi, p, T_u, f)$ to be smaller than or equal to 1.

$$\begin{aligned}
 F_1(\phi, T_u, p, f) = & c_1 + c_2\phi + c_3\frac{T_u}{T_0} + c_4\frac{p}{p_0} + c_5f + c_6\phi^2 \\
 & + c_7\left(\frac{T_u}{T_0}\right)^2 + c_8f^2 + c_9\phi\left(\frac{T_u}{T_0}\right) + c_{10}\phi\left(\frac{p}{p_0}\right) \\
 & + c_{11}\phi f + c_{12}\left(\frac{T_u}{T_0}\right)f + c_{13}\phi^3 + c_{14}f^3 \\
 & + c_{15}\left(\frac{T_u}{T_0}\right)\phi^2 + c_{16}\left(\frac{p}{p_0}\right)\phi^2 + c_{17}\phi\left(\frac{T_u}{T_0}\right)^2 \\
 & + c_{18}f\left(\frac{T_u}{T_0}\right)^2 + c_{19}\phi f^2 + c_{20}\left(\frac{T_u}{T_0}\right)f^2
 \end{aligned} \quad (5)$$

$$F(\phi, T_u, p, f) = \min[1, F_1(\phi, T_u, p, f)] \quad (6)$$

The quality of the fit was checked by comparing the predicted u_l values against the fitted data points and a batch of test data. This batch consisted of calculated u_l data that were discarded in the fitting process at various equivalence ratios, pressures and unburned mixture temperatures. Table 6 provides an overview of the quality of the fit for ethanol–air mixtures, listing the average relative residual (Eq. (7)), average absolute relative residual (Eq. (8)), minimum and maximum residual and the percentage of data points that are predicted to within $\pm 10\%$ and $\pm 20\%$. As can be seen the relationship captures 93% of the fitted data to within $\pm 20\%$ and has a mean absolute relative residual of 7.38%. For the test data it has a mean absolute relative residual of 8.58% and captures 88.77% to within $\pm 20\%$.

$$\text{average relative residual} = \frac{1}{N} \sum \left(\frac{u_{l,\text{pred}} - u_l}{u_l} \right) \quad (7)$$

$$\text{average absolute relative residual} = \frac{1}{N} \sum \left(\frac{|u_{l,\text{pred}} - u_l|}{u_l} \right) \quad (8)$$

It is important to note that the largest differences occur for extreme values of ϕ (0.5, 2.0), p (≤ 5 bar, ≥ 85 bar) which is a result of the polynomial form of the equations. Additionally, the calculations showed aberrant behavior at the highest temperatures (≥ 800 K) due to self-ignition. This led to a more than exponential rise of u_l in terms of unburned mixture temperature and burning

Table 6
Fitting statistics for Eq. (2) (full fit, $f \geq 0$ vol.%).

	$u_{l,\text{pred}}$ fitted data (%)	$u_{l,\text{pred}}$ test data (%)
Average rel. residual	0.61	-2.74
Average abs. rel. residual	7.38	8.58
Maximum residual	69.72	40.43
Minimum residual	-81.41	-60.82
Data within ± 10	73.64	67.95
Data within ± 20	93.30	88.77

velocities that increased with rising diluent fraction. These results were not retained to fit the equations, but were included in this quality check.

Having determined the coefficients for the correction term F , the full correlation is now known. It consists of Eq. (2) whereby $\alpha(\phi, p)$, $u_{l0}(\phi, p)$ and $F(\phi, T_u, p, f)$ are computed through Eqs. (3)–(6) respectively, making use of the coefficients listed in Tables 4 and 5, p_0 and T_0 as given above, f the volume fraction of residuals and u_l given in cm/s. A C++ implementation of the correlation is available online as supplementary material with this paper.

4. Flame thickness correlation

4.1. Evaluation of published correlations

Flame thickness influences the combustion process through its defining effect on flame–turbulence interaction. Thicker flames are less sensitive to small eddies and will therefore experience lower degrees of turbulence induced flame wrinkling. Some recent turbulent combustion models incorporate the effects of flame thickness, requiring an accurate knowledge of this quantity at engine-relevant conditions. For example, Bougrine et al. [32] used an efficiency function $\Gamma(u'/u_l, l_t/\delta_l)$ proposed by Charlette et al. [6], which measures the ability of the different turbulent scales to wrinkle the flame front. Where u' is the intensity of velocity fluctuations, l_t is the turbulence integral length scale and δ_l is the laminar flame thickness.

The model was derived by Charlette et al. from DNS of interactions between single vortices and a flame in order to measure the effective strain-rate of vortices of different characteristic sizes and speeds. Based on these DNS results they fitted an effective flame wrinkling function Γ that accounts for all scales relevant to engine combustion. This efficiency function evaluates the surface-producing flame stretch across different combustion regimes. Obviously the influence of flame thickness is most pronounced for the thickened wrinkled flames regime ($Ka > 1$) and the thickened flame regime ($Da < 1$). In these regimes, the thickness of the flame limits the smallest turbulence scales in wrinkling the flame front. In the wrinkled flame regime the assumption of infinitely thin flamelets is more valid and the influence of flame thickness is negligible.

To limit computational efforts in engine simulations, this laminar flame thickness δ_l is also determined using a correlation. For conventional hydrocarbon fuels different flame thickness correlations have been proposed in the literature. The Zel'dovich correlation expresses the flame thickness in terms of the laminar flame speed and the fresh gases properties [33]:

$$\delta_l^{\text{Zeldovich}} = \left(\frac{\lambda}{\rho \cdot C_p} \right)_u \frac{1}{u_l} \quad (9)$$

where λ is the thermal conductivity, ρ the density and C_p the heat capacity of the fresh gases. Another correlation was proposed by Blint [33] including a correction for burned gases properties using a Sutherland law for λ and a constant Prandtl number of 0.7:

$$\delta_l^{\text{Blint}} = 2 \cdot \left(\frac{\lambda}{\rho \cdot C_p} \right)_u \frac{1}{u_l} \cdot \left(\frac{T_b}{T_u} \right)^{0.7} \quad (10)$$

where T_b is the burned gases temperature. We compared these correlations against our database of calculated flames, where the flame thickness was estimated from the temperature gradients between the fresh and burned gases zones:

$$\delta_l^{\text{CHEM1D}} = \frac{T_b - T_u}{\max(dT/dx)} \quad (11)$$

Several flame thickness definitions have been proposed in the literature including definitions based on the temperature gradients, characteristic chemical time and the heat release rate.

Blint states that a flame width definition based on temperature gradient is the most rational selection to provide an unambiguous specification of the laminar length scale, since it can be directly determined from the temperature profile and incorporates both transport and heat release [33]. Still, this definition is not optimal as it implicitly assumes that temperature profiles have the same shape for all conditions, which might not necessarily be the case, especially around flammability limits [32].

Fig. 6 compares the flame thickness calculated using the mechanism of Li et al. [21] against the correlations of Zel'dovich and Blint for varying equivalence ratio and at several pressures ($T_u = 700$ K, $f = 0$ mol%). In Fig. 7 the correlations are compared against calculation results for varying diluent ratio and at different unburned mixture temperatures ($p = 15$ bar). Also included in the figures is a new correlation, which is discussed in Section 4.2. The Zel'dovich correlation underestimates the calculated flame thicknesses at all conditions. Blint's correlation shows better agreement, but errors increase for lean and rich mixtures at low pressures and highly diluted mixtures at low temperatures. In these conditions the largest overestimations of calculated flame thicknesses can be seen, both in absolute values as in percentual differences.

The same behavior was noted by Bougrine et al. when they compared calculated flame thickness for methane–air flames against the correlations of Zel'dovich and Blint [32]. Bougrine et al. finally retained the correlation of Blint in their modeling work, because it predicts the right trends for flame thickness with pressure, unburned mixture temperature and equivalence ratios. Moreover the largest errors are seen in conditions that are of lower importance to engine modeling work and where the validity of the flame thickness definition based on temperature gradients is questionable. In this study, however, it was noted that the use of Blint's

Table 7

Comparison of δ_l predicted by Blint's correlation (Eq. (10)) against calculation results using the ethanol oxidation mechanism of Li et al. [21].

	δ_l^{Blint} (%)
Average rel. residual	13.45
Average abs. rel. residual	17.27
Maximum residual	46.23
Minimum residual	−81.76
Data within $\pm 10\%$	30.88
Data within $\pm 20\%$	59.16
Data within $\pm 30\%$	81.52

correlation led to an average overestimation of the calculated laminar flame thickness of over 13%. Also, it was observed that in less than 31% of the calculated cases, the flame thickness was predicted within 10% (see Table 7). For this reason we attempted to make slight adjustments to Blint's correlation in order to improve its predictive performance.

4.2. New laminar flame thickness correlation

A database of over 1500 calculated flames in the range of 1–85 bar for pressure, 400–900 K for the fresh gas temperature, 0.5–2 for ϕ and diluent ratios up to 50 vol.% was used to evaluate the Blint correlation for ethanol–air mixtures (see Table 7). The same database can now be used to fit an improved flame thickness correlation.

Knop [34] adapted Blint's correlation to hydrogen combustion by adding a multiplying factor κ . This factor κ is unity for conventional hydrocarbons and depends on the equivalence ratio ϕ , fresh gas temperature T_u and the pressure p for hydrogen:

$$\kappa = \alpha \cdot f(\phi) \cdot g(T_u) \cdot h(p) \quad (12)$$

where $\alpha = 3.37$ and f , g and h are second order polynomial functions of ϕ , T_u and p respectively. In engine-like conditions κ ranged from 2 to 7, which means that for the same laminar burning velocity the hydrogen flame is clearly thicker than the hydrocarbon counterpart.

For ethanol–air flames the errors associated with the Blint correlation were limited to 30% at over 80% of the calculated conditions (see Table 7). Consequently, it was preferred to keep the form of the correlation (Eq. (10)), add a multiplying constant κ and change the exponents for u_l (from -1 to β) and T_b/T_u (from 0.7 to 0.5).

The Sutherland law for λ in its original form employs an exponent of 0.5, so this value was used instead of 0.7 for the correction for burned gases properties:

$$\lambda_b = \left(\frac{T_b}{T_u} \right)^{0.5} \cdot \lambda_u \quad (13)$$

The proposed correlation for laminar flame thickness of ethanol–air mixtures is given by Eq. (14):

$$\delta_l^{\text{New}} = \kappa \cdot \left(\frac{\lambda}{\rho \cdot C_p} \right)_u \cdot u_l^\beta \cdot \left(\frac{T_b}{T_u} \right)^{0.5} \quad (14)$$

where κ and β are constants that were determined by minimizing the SSD between the flame thicknesses in our flame database and values predicted by the correlation. Two sets of constants were computed (see Table 8). The first set was fitted using the laminar burning velocity values calculated by CHEM1D as an input for the correlation. The second set employed the correlation proposed in the first part of this paper to predict u_l . Note that u_l is expressed in cm/s, while the correlation gives δ_l in cm. The predictive performance of Eq. (14) using both sets of values for κ and β is summarized in Table 8. A first

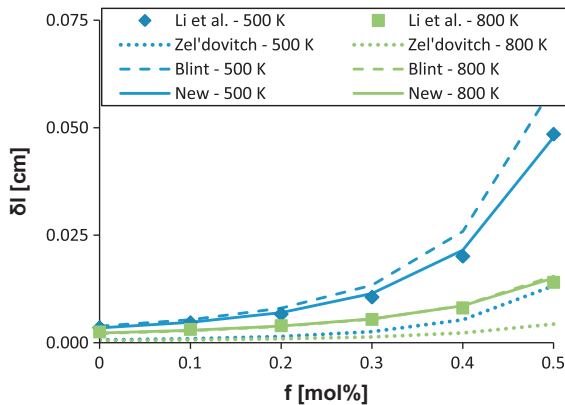


Fig. 6. Laminar flame thickness variations with the equivalence ratio ϕ at different pressures ($T_u = 700$ K, $f = 0$ mol%).

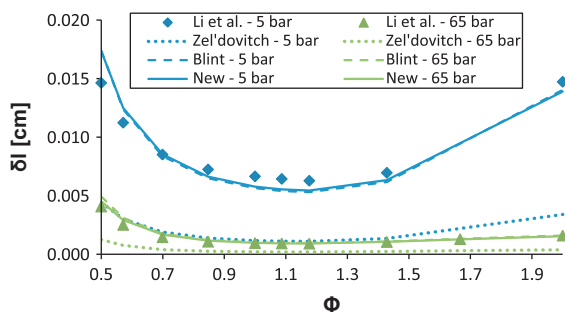


Fig. 7. Laminar flame thickness variations with diluent ratio f at unburned mixture temperatures ($p = 15$ bar, $\phi = 1$).

Table 8

Comparison of δ_l [cm] predicted by the adapted Blint correlation (Eq. (10)) against calculation results. The left column summarizes the quality of the fit using calculated u_l [cm/s] values (u_l^{CHEM1D}), the right column for the fit using u_l [cm/s] values predicted by Eqs. (2)–(6) ($u_l^{\text{corr.}}$).

	$\delta_l(u_l^{\text{CHEM1D}})$	$\delta_l(u_l^{\text{corr.}})$
κ	0.9505	0.9484
β	−0.9346	−0.9382
Average rel. residual (%)	1.12	0.75
Average abs. rel. residual (%)	6.63	8.48
Maximum residual (%)	25.70	52.71
Minimum residual (%)	−81.68	−56.49
Data within $\pm 10\%$ (%)	78.43	70.65
Data within $\pm 20\%$ (%)	96.07	91.30
Data within $\pm 30\%$ (%)	98.75	96.00

thing to notice is that both sets are almost identical, which confirms the predictive performance of our laminar burning velocity correlation. The correlation using the CHEM1D results predicts the calculated flame thickness to within 10% in 78% of the observed conditions. In almost all conditions the errors on δ_l are below 20%. The good fit with calculation results can also be observed in Figs. 6 and 7 ('New' correlation). The largest deviations occur in conditions that are on the edge of self-ignition (high pressures and temperatures) or flame extinction (very rich or lean, high amounts of residual gases). In these conditions the results of the chemical kinetic calculations and the definition of flame thickness based on temperature gradients are questionable.

The flame thickness correlation using our own correlation for burning velocity performs somewhat worse with only 70% of the calculated flame thickness being predicted to within 10%. This is caused by an underprediction of the laminar burning velocity at low pressures (≤ 5 bar). This results in an overestimation of the associated flame thickness. These low pressure conditions, however, are of minor importance for engine simulation purposes.

5. Conclusions

Bio-ethanol is an interesting alternative fuel for use in spark-ignition engines. The use of an engine cycle code valid for this light alcohol, could help to explore its full potential. Important building blocks for most predictive engine codes are the laminar burning velocity and flame thickness of the fuel–air–residuals mixture at instantaneous pressure and temperature. These parameters are conveniently implemented in engine codes by using correlations that give their values in terms of pressure, temperature and composition of the unburned mixture.

In the first section of this paper, it was shown that for ethanol there is a lack of burning velocity correlations suitable for use in engine codes. In fact experimental laminar burning velocity data at engine-like conditions is very scarce, especially data at elevated pressures and for diluted mixtures. Moreover, most of the published data and correlations are compromised by the effects of flame stretch and the occurrence of flame instabilities.

Computationally, these effects can be avoided by calculating one-dimensional, planar adiabatic flames using chemical oxidation mechanisms. A number of contemporary chemical mechanisms for the oxidation kinetics of ethanol were selected from literature. The results from calculations with these mechanisms were compared against reliable experimental data for the laminar burning velocity for a range of pressures, temperatures, equivalence ratios and diluent ratios. Based on these studies the mechanism of Li et al. [21] was retained to calculate laminar flames for a wide range of engine-like equivalence ratios (0.5–2), pressures (5–85 bar), unburned mixture temperatures (400–900 K) and diluents fractions (0–50 vol.%). It was noted that at temperatures above 900 K

the mixture self-ignited, which renders the calculation results useless.

Using this database of computed flames, we constructed new correlations for the laminar burning velocity and flame thickness of ethanol–air mixtures. For the laminar burning velocity, it was shown that existing correlation forms cannot capture the strong interaction effects between equivalence ratio, pressure, temperature and diluent fraction. Therefore, a new correlation form was proposed:

$$u_l(\phi, p, T_u, f) = u_{l0}(\phi, p) \left(\frac{T_u}{T_0} \right)^{\alpha(\phi, p)} F(\phi, p, T_u, f)$$

where $\alpha(\phi, p)$, $u_{l0}(\phi, p)$ are third order polynomial functions of ϕ and p with various cross-terms. The influence of residuals on the burning velocity is incorporated in a separate correction term $F(\phi, T_u, p, f)$, which is also a polynomial function of ϕ , T_u , p and diluent fraction f . The proposed correlation form closely fits the detailed chemical kinetics results.

An evaluation of different flame thickness correlations demonstrated that the correlation of Blint predicts the right trends for δ_l of ethanol in terms of ϕ , T_u , p and diluent fraction. Still, some constants in the correlation were slightly adapted in order to better match our calculation results. Flame thicknesses were estimated from the temperature gradients in the computed flames.

The developed correlations can now be implemented in an engine cycle code. Future work will focus on further validating the correlations by comparing them to our own experimental laminar burning velocity values and pressure traces obtained during engine experiments.

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