Simultaneous multispectral imaging of flame species using Frequency Recognition Algorithm for Multiple Exposures (FRAME)

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ABSTRACT

Imaging the interaction between different combustion species under turbulent flame conditions requires methods that both are extremely fast and provide means to spectrally separate different signals. Current experimental solutions to achieve this often rely on using several cameras that are time-gated and/or equipped with different spectral filters. In this work we explore a technique called Frequency Recognition Algorithm for Multiple Exposures (FRAME) as an alternative solution for instantaneous multispectral imaging of flame species. The method is based on exciting different species with different spatial “codes” and to separate each signal component using a spatial frequency-sensitive lock-in algorithm. This methodology permits the signal from several different species to be recorded at the exact same time with a single camera. Furthermore, since the signals are recognized based on the superimposed spatial codes, there is no need for spectral separation prior to detection. The entire fluorescence envelope from each species can thus, in principle, be detected. In the current work, we present simultaneous planar laser-induced fluorescence imaging of OH and CH₃O in a turbulent dimethyl ether (DME)/air flame.

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In this paper, we explore an alternative method to tackle the problem associated with multispectral data acquisitions. The method, which is referred to as Frequency Recognition Algorithm for Multiple Exposures (FRAME) in the literature [23,24], employs a coded laser-probe illumination strategy to distinguish between different signal components. Dorozynska and Kristensson recently demonstrated how this coding strategy could be exploited in order to acquire multispectral 2D information instantaneously [25]. In the current study, we continue exploring the potential of the FRAME technique for simultaneous multispectral imaging, primarily seeking to investigate the merits of FRAME for gas-phase, combustion diagnostics. To reach sufficient signal-to-noise (SNR) levels we transfer the wide-field FRAME imaging concept adopted by Dorozynska and Kristensson into a laser sheet arrangement. With this setup we here demonstrate, for the first time, the ability to perform simultaneous multispectral PLIF imaging of combustion species – OH and CH$_2$O – with a single camera by means of FRAME.

2. Frequency Recognition Algorithm for Multiple Exposures (FRAME)

FRAME is an imaging technique that allows a mono-camera setup to record several laser-probe images in one single acquisition [23–25]. Under normal circumstances such an approach would yield an image wherein all the different 2D signal components would be irreversibly mixed. Separating the different images would require knowledge about the spatial distribution and the concentration of the sample constituents – the quantities one seeks to determine. FRAME solves this “problem” by employing a coded illumination strategy, where each individual fluorescence image becomes encoded with a specific periodic line structure (pure sinusoidal or top-hat). The illumination scheme is similar to that used for Structured Illumination Planar Imaging (SLIPI) [26–28] but unlike SLIPI, FRAME accesses all data from a single recording, albeit at the cost of a reduced spatial resolution. The 2D signal from each laser-probe can thus be distinguished and separated from each other in the data post-processing by means of a frequency-sensitive spatial lock-in algorithm [29,30]. Here we will first describe the FRAME principle conceptually followed by a mathematical description of the method and the lock-in algorithm.

In the conceptual example presented in Fig. 1, a sample with three different constituents is illuminated using the coded illumination approach. In Fig. 1a, the sample is illuminated with a single modulated (frequency $\nu_1$) laser-probe at a wavelength of $\lambda_1$, sensing the spatial distribution of one constituent. The sample structures and laser light are superposed multiplicatively, effectively creating three “copies” of the sample structures in reciprocal space (Fourier domain), one in the origin and two centered at $\pm \nu_1$. If the modulation frequency is sufficiently high, the “copies” will be

![Image](image-url)

**Fig. 1.** A conceptual illustration of the FRAME principle. A sample with three different constituents is illuminated with differently coded light fields. (a) A single modulated laser-probe illuminate the sample, creating two “copies” of the sample structures of one constituent at $\pm \nu_1$. (b, c) By illuminating the sample with differently coded laser-probes, it is possible to transfer data from the different constituents into different regions in reciprocal space.
isolated from each other, as illustrated in Fig. 1a. In Fig. 1b and c differently coded laser sources excite the remaining constituents of the sample simultaneously, thus transferring “copies” of the different sample structures into different, isolated regions in reciprocal space. The acquired image – containing data from all constituents – is then analyzed with an algorithm that only extracts information that is modulated with a certain spatial frequency.

To explain the FRAME principle and lock-in algorithm mathematically, consider the case given in Fig. 1b, where a sample is illuminated with two differently coded laser-probes. The signal recorded by the camera can be expressed as follows:

\[
I(x, y) = S_1(x, y)(1 + \sin(2\pi v_1 x + 2\pi v_1 y)) + S_2(x, y)(1 + \sin(2\pi v_2 x + 2\pi v_2 y))
\]

(1)

where \(I(x, y)\) is the acquired image, \(S_1\) and \(S_2\) are the two sample responses and \(v\) are spatial frequencies (the spatial phase of the modulation is omitted for reasons of clarity). From this data we now aim to extract the sample responses individually, here derived for the extraction of \(S_1\). Multiplying Eq. (1) with a sine wave having the modulation frequency \(v = v_{1,x} + v_{1,y}\) yields:

\[
I(x, y) \sin(k v_{1,x} x + k v_{1,y} y) = \frac{1}{2} \left( S_2 \cos(k(v_{1,x} - v_{2,x})x + 2\pi(v_{1,y} - v_{2,y})y) - S_2 \cos(k(v_{1,x} + v_{2,x})x + k(v_{1,y} + v_{2,y})y) + 2S_1 \sin(k v_{1,x} x + k v_{1,y} y) - S_1 \cos(2k v_{1,x} x + 2k v_{1,y} y) + 2S_2 \sin(k v_{1,x} x + k v_{1,y} y) + S_1 \right)
\]

(2)

Frequency analysis of Eq. (2) (where \(k = 2\pi\)) reveals several different frequency components, yet only a single DC component – the last \(S_1\) term. Thus, by applying a low-pass filter with a cut-off frequency \(v_c\), where \(v_c < v_{1,x} + v_{1,y}\) and \(v_c < v_{2,x} + v_{2,y}\), all terms but the \(S_1\) term are rejected, thus revealing the sample structures of \(S_1\). The second sample constituent, \(S_2\), is then accessed in a similar fashion, i.e. by multiplying Eq. (1) with a sine wave having the modulation frequency \(v = v_{2,x} + v_{2,y}\). It should be noted that to avoid ambiguities caused by the spatial phases, which were omitted in the derivation, Eq. (1) also needs to be multiplied with \(\cos(k v_{1,x} x + k v_{1,y} y)\) Combining both expressions cancels out the spatial phase, see also Refs. [29,30] for more detailed derivations of the spatial lock-in algorithm (albeit in one dimension only).

Figure 2 illustrates the mathematical procedure of the lock-in algorithm graphically. Figure 2a shows an unprocessed image containing planar laser-induced fluorescence signals from both OH and CH\(_2\)O, generated using the FRAME setup used in the current study. The Fourier transform of Fig. 2a (Fig. 2b) reveals the two image components stored simultaneously on chip. Multiplying the input image with a 2D sine wave that exactly matches the modulation frequency of the OH signal effectively demodulates the fluorescence from OH (Fig. 2c), i.e. the complex information is shifted so that the modulation frequency of OH is placed at the origin. The rearranged dataset is then filtered using a low-pass Gaussian filter (see inset), thus extracting the OH image and removing the information arising from CH\(_2\)O. Calculating the inverse Fourier transform turns spatial frequencies into intensity values, revealing the OH distribution (Fig. 2d). The procedure is then repeated for the fluorescence signal emitted by the CH\(_2\)O molecule (Fig. 2e).
Fig. 3. FRAME experimental setup. Two identical optical arrangements were created for the laser excitation sources. In the setup, a transmission grating was illuminated and a cylindrical lens was used to form an image of the grating. A spatial filter, placed in the Fourier plane of the first cylindrical lens, rejected all but the ±1 diffraction orders of the grating in order to create a pure sinusoidal intensity modulation (by interference). A second cylindrical lens was then used to compress the light field into a thin sheet. Two mirrors were used to alter the angle of incidence, to transfer the fluorescence signal from each laser sheet into different regions in the Fourier domain of the acquired image. An ICCD camera (Princeton Pi-MAX4) recorded the fluorescence emitted at 90°.

Fig. 4. Simultaneously acquired multispectral images of OH and CH$_3$O in a Bunsen flame. (Top row) Four unprocessed images, as seen by the camera, recorded using the FRAME setup. (Bottom row) Multispectral information after analyzing each image with the lock-in algorithm (filter size $\sigma = 0.17$).

The two images are false colored and merged into a multispectral image.

3. Experimental setup

Figure 3 shows a schematic of the optical arrangement used in the experiment. A frequency-tripled Nd:YAG laser ($\lambda = 355$ nm, pulse energy $\approx 100$ mJ) was used to excite the formaldehyde molecules, while the hydroxyl molecules were stimulated using a frequency doubled dye laser ($\lambda = 283$ nm, pulse energy $\approx 10$ mJ), pumped by a pulsed Nd:YAG laser ($\lambda = 532$ nm). The optical arrangements were identical for the two different light sources. After being expanded and collimated, the beams were each guided through a Ronchi grating (10 lp/mm), diffracting the beams into several different diffraction orders. To form the sinusoidal line pattern in each beam path, a cylindrical lens ($f = 150$ mm) was used in conjunction with a spatial filter, which only permitted the ±1 orders of diffraction. Employing a spatial filter is not strictly required – the grating can be imaged directly onto the measurement region – but it does simplify the alignment procedure since the exact distance between the cylindrical lens and the sample becomes less crucial (the two orders create the intensity modula-

One drawback with this approach is the reduction of the laser energy, where $\sim 50\%$ is lost due to reflection upon the Ronchi grating and thereafter $\sim 70\%$ of the remaining energy is blocked by the spatial filter. Although the final photon flux was sufficient for the current proof-of-concept investigation, applications with low SNR conditions may require alternative optical configurations where the laser power is better preserved. Kristensson et al. employed a structured illumination scheme based on a double-wedged glass plate, which yields insignificant losses in photon flux [31]. Another potential solution would be to create the intensity modulation by means of a phase grating – a transparent optical component that diffracts the transmitted light directly into the desired ±1 orders. Employing such lossless structured illumination schemes thus have the potential of improving the photon flux by a factor of $\sim 6$, although this needs to be confirmed through experimental work.

A second cylindrical lens ($f = 1000$ mm) was then used to compress each laser beam into a thin sheet of light ($\sim 200 \mu$m thickness). Maximizing the modulation depth for each beam path requires fine-tuning of the orientation of the grating as well as both cylindrical lenses. Real-time monitoring of the Fourier transform of the acquired image can thus facilitate in this procedure. Finally,
two mirrors were used in both beam paths to guide the laser sheets onto the sample from different angles in order to transfer the corresponding fluorescence into different regions in the Fourier domain. An intensified CCD (ICCD) with a pixel resolution of 1024 × 1024, positioned at a 90° angle configuration, was used to image the fluorescence from the two species simultaneously using a Halle lens (local length = 100 mm, f_p = 2). A 290 nm long-pass filter and 355 nm notch filter were applied to remove the scattering signals. The burner was a Bunsen type burner, providing a pre-mixed, slightly fuel rich DME/air flame.

4. Results and discussion
4.1. Multispectral imaging results

Figure 4 shows four examples of multispectral images, acquired simultaneously using the planar FRAME setup. The top row in Fig. 4 shows the raw data, as seen by the camera. The magnified regions highlight how the signal from OH and CH_3O differs in terms of their modulated structure, making it possible to, even by eye, distinguish the signal from the two species. Each set of data is then analyzed using the FRAME algorithm, which separates the species, the result of which is shown in the bottom row (false coloring). Note that no correction for e.g. flat-field compensation or background suppression has been applied to the data. Since the data are not quantitative, the signal from each species is histogram-adjusted and normalized to unity in the presented data. The data show that the formaldehyde is formed in the area where the pre-heat zone and the reaction zone meet whereas the OH molecule is present in the product zone, which is in agreement with past knowledge [13,15,32–34]. The formaldehyde visible in the high temperature regions is due to the relatively low SNR for the formaldehyde signal, which, as mentioned above, is primarily due to the losses caused by the approach currently used to generate the line structures. The use of e.g. phase gratings or double-wedged glass plates should increase the SNR for improved sensitivity.

Figure 5 presents statistical analysis performed on the acquired dataset (200 snapshots), showing averaged images, standard deviations in 2D and mode analysis (two most dominant modes). Mode analysis (or Proper Orthogonal Decomposition, POD) is an analytical tool to statistically investigate a dynamic behavior (in particular oscillatory modes) and is based on decomposing an ensemble of images (or 1D arrays) into a set of Eigen functions (modes). The modes represent fluctuations from the mean field, where any snapshot in the dataset can be reconstructed using a linear combination of the modes. For more details on the theory of POD and its applications for combustion studies, see Refs. [35–38].

In 1998, Paul and Najm proposed that taking the pixel-by-pixel product of OH and CH_3O images yields an image that displays the reaction rate [33]. The ability to collect both these data simultaneously as well as an accurate pixel-by-pixel correlation is thus key for such data analysis. Figure 6 shows four typical FRAME measurements using the current setup, from which this product is calculated. Since both images are acquired using the exact same collection optics and camera system, corrections procedures to reduce measurement uncertainties associated with these parameters are not needed. Note that the relatively low SNR of the CH_3O leads to both the non-zero values of reaction rate in the high temperature regions as well as the zero values of heat release in the reaction zone.

4.2. Spectral filtering

The ability to access multispectral information simultaneously with a single detector also opens up for new measurement possibilities for e.g. spectral filtering and suppression of interferences. For example, detecting species having fluorescence signals that partly or entirely overlap spectrally is technically challenging with multispectral imaging schemes that rely on spectral filters. Examples of combinations of combustion species whose fluorescence signals are difficult to separate spectrally are (1) acetone, CH and CH_3O [39–45] and (2) OH and toluene [41,46–48]. Using several time-gated cameras to acquire the signals sequentially solves the problem, but it is an expensive solution. Since FRAME uses spatial codes to distinguish between signal components, it could, in principle, be used to detect and separate species with similar – or even identical – spectral emission characteristics. Note that notch filters may still be required to prevent detection of laser scattering. A further potential advantage of avoiding spectral filters to separate signal components by means of FRAME is an improved SNR, since it may allow detection of the entire fluorescence envelope rather than only selecting (spectrally) the non-overlapping part. Of course, this would require a more light-efficient FRAME setup, as those mentioned in the Experimental setup section, so that the lower excitation flux does not negate the improved emission collection.

Another potential benefit with FRAME concerns the detection and suppression of interfering signal components arising from broadband absorbers and/or stray light generated by e.g.

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**Fig. 5.** Statistical analysis of the acquired dataset (200 snapshots) for both OH and CH_3O. The figure shows mean fields, standard deviations and the two most dominant modes from POD analysis for both combustion species.
Figure 6. Pixel-by-pixel product of OH and CH$_2$O images. Taking the product of the OH (top row) and the CH$_2$O (middle row) images yields intensity maps that are closely related to the reaction rate (bottom row) [15,31].

4.3. Spatial resolution

The ability to store several images simultaneously on the camera sensor comes at the price of a reduced spatial resolution per extracted image. For traditional photography, the spatial resolution achievable depends mainly on (1) the number of pixels of the detector and (2) the modulation transfer function (MTF) of the imaging system, i.e. the optical properties. However, laser sheet imaging in general, and planar laser-induced fluorescence imaging of gaseous samples in particular, has one more factor that influences the spatial resolution – the thickness of the laser sheet. This can be understood by considering that the sample extends in 3D and that the laser sheet is generating a signal in the entire volume it is encountering. Although, it is also important to realize that not all objects carry structural details finer than those resolvable by a given imaging system. In these cases, the data is over-sampled.

The spatial resolution of images extracted from a FRAME measurement depends on two additional factors; the spatial frequency of the superimposed intensity modulation and the low-pass filter used in the spatial lock-in algorithm. These two parameters are closely linked. Increasing the size of the low-pass filter means including more spatial frequencies, which yields an improved spatial resolution. However, the size of the filter cannot exceed a certain limit, after which cross-talk between neighboring images (in reciprocal space) becomes apparent (see example in Fig. 7). Generally, the more isolated each image copy is in the Fourier domain, the better the end resolution becomes. It is therefore advisable to maximally disperse the image copies in reciprocal space and use the highest intensity modulation frequency resolvable by the imaging system. Unfortunately, intensified (multi-channel plate, MCP) cameras tend to have an MTF that rolls off relatively fast, which is because fine structural details are lost in the conversion from photons to electrons and then back to photons again. This characteristic becomes a limiting factor when employing the FRAME concept with intensified cameras, because the intensity modulation frequency must be kept relatively low (to be kept within the borders of the MTF). Consequently, this affects the settings of the low-pass filter whose dimensions also become relatively small.

Figure 7 illustrates how the spatial resolution of the extracted images is influenced by the settings of the low-pass filter. The example shows a raw, unprocessed image that contains both OH and CH$_2$O data together with its Fourier transformation after the OH image copy has been demodulated (compare with Fig. 2c). The OH image is then extracted using five differently sized filters, where the $\sigma$-value in each frame indicates the integrated area of the low-pass filter (acceptance of spatial frequencies) given as a percentage

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of the area of the Fourier domain. A filter that covers the entire reciprocal space has a $\sigma$-value of 100%. At low values of the filter, most image details are lost. Although, already at 0.06% the OH distribution starts to become apparent. When the filter is increased beyond a certain limit information from neighboring images (or the 0th order) fall within the filter and creates an interfering signal. The last example in Fig. 7 ($\sigma = 0.39\%$) shows such interference, which is manifested as lines. Note that in this particular example, the cross-talk comes from the 0th order data (originally contained in the origin) and not the neighboring CH$_2$O image copy.

The FRAME concept relies on the fact that the power spectrum of an image is strongly shifted towards the low spatial frequencies [51]. To further illustrate this fact, we present a set of synthetic images of a resolution target that have been processed with different filter settings (Fig. 8). The figure shows results from three of the filters in Fig. 7 where, as before, $\sigma$ indicates the integrated area of the low-pass filter divided by the full area of the sensor, given in percentage. Also included in Fig. 8 is an image of the resolution target that shows the spatial resolution achievable with a 200$\mu$m thick laser sheet setup. Note that the calculations are based on a detection system that has a pixel resolution of 1024 $\times$ 1024 and that FRAME would benefit, both in terms of image-storing capabiliities as well as the spatial resolution, of having a higher number of pixels (as is true for nearly all types of imaging measurements). This type of analysis can act as guidance, making it possible to determine how a FRAME setup should be arranged depending on what structures one desires to observe. In the current experiment and with the current imaging conditions, the maximum size of the low-pass filter was 0.17%, which gives a spatial resolution of $\sim 0.4$ mm (2.3 lp/mm), i.e. corresponding to the resolution achievable with a 0.4 mm thick laser sheet. This spatial resolution was deemed sufficient for this proof-of-concept demonstration of FRAME, allowing observation of the spatial distribution of both the OH- and CH$_2$O molecules on a macroscopic scale. However, more detailed studies of e.g. the overlap between different combustion species using FRAME would require an improved spatial resolution to avoid ambiguities caused by spatial broadening. Observing finer details with FRAME can be accomplished in two ways. First, by increasing the intensity modulation frequency a larger low-pass filter can be used in the FRAME image extraction process. For example, increasing this frequency by a factor of $\sim 1.5$ would permit the use of $\sim 6$ times larger filter. This calculation illustrates an important feature; the relationship between filter size and modulation frequency is not linear and a relatively small increase in modulation frequency can yield a significant improvement in spatial resolution. Second, reducing the field-of-view (currently $\sim 25 \times 25$ mm) through optical magnification also improves the spatial resolution, given that the modulation frequency is increased accordingly (achieved by e.g. down-collimating the excitation sheets).

Figure 9 illustrates the relationship between modulation frequency and filter size on a set of synthetically generated images.
A very low modulation frequency, case (a), leads to a condition where the coded information cannot be accessed due to a too large overlap with the low frequency data. As the modulation frequency is increased the overlap between structures in the reciprocal space reduces, which allows access to the frequency-shifted data. Notice how a higher modulation frequency yields conditions that permit larger filters without introducing cross-talk with lower spatial frequencies (circles in Fig. 9).

The spatial resolution required for drawing conclusions based on visual observations or statistic analysis is highly case-dependent. Mode analysis (POD), which uses a large dataset to extract Eigen functions (or spatial modes), is an powerful analytical tool that FRAME imaging potentially is well-suited for since it requires snapshot data. To investigate how sensitive POD is to spatial resolution, the acquired dataset of 200 snapshot images were analyzed using the FRAME algorithm with 13 different filter sizes (σ) for both the OH and CH₂O signals. Mode analysis was then performed on all 13 datasets for both combustion species, and the two most dominant modes for each individual case were extracted. To determine how the spatial resolution of the images that are extracted by the FRAME algorithm is affecting the set of Eigen modes, each set of 13 modes was compared with its corresponding reference case. The reference case was extracted using a filter size of σ = 0.17 which was the largest filter size used in this current
analysis (see also Fig. 7). The comparison between the two most dominant modes in the reference case (denoted $M_{j}(\sigma, 0.17)$ in Fig. 10, where $j$ = mode number) and the remaining 12 sets of mode images was based on computing their scalar product, giving values between zero and unity. A value of one means that the mode analysis is unaffected by the reduction of the filter size, and thus the same modes can represent both image sets. If, however, the scalar product is zero there is no correlation between the mode image and its reference case (image vectors are orthogonal). The results from this investigation are displayed in Fig. 10. A filter size, $\sigma$, above $\sim0.12$ yields virtually no effect (scalar product of $\geq 0.99$) on the POD analysis on the set of CH$_2$O data, whereas the equivalent number for the set of OH data is $\sigma \approx 0.06$. In practice, this means that the same set of Eigen modes can be used to represent OH and CH$_2$O data if the filter size is kept above these numbers. However, the Eigen-mode analysis starts to be affected if smaller filter widths are used. Although the results are slightly different for OH and formaldehyde – which we attribute to differences in their spatial distribution – the results show that the loss in image resolution caused by FRAME does not have a significant impact on mode analysis.

4.4. Beam steering

Refraction occurring due to a sudden change in refractive index – beam steering – is a potential concern for FRAME imaging since the method is based on extracting light that carries a specific periodic line structure. Significant beam steering occurring in the flame will thus distort the periodic pattern, which would lead to a reduced signal level. For laser sheet imaging based on structured intensity profiles, beam steering will generate a shift in the line structure that, in turn, will lead to an asymmetry in the extracted species image, with a stronger signal on the entrance side (analogous to laser extinction). To investigate whether effects of beam steering can be observed in the data, the excitation line structures are traced across the field-of-view, see Fig. 11. Focusing in particular on regions before and after the reaction zone where the temperature gradient is the steepest and thus the impact of beam steering should be the greatest. In the analysis, each modulated line is treated as a single beam and any departure from its optical path should be visually apparent in the recorded data. The analysis shows, however, no visible traces of beam steering, which is most likely due to the relatively short distance light travels between the two borders of the reaction zone. Measurement issues associated with beam steering may, however, be more pronounced when probing larger combustion systems using FRAME.

5. Summary

In summary, we have, for the first time, applied the FRAME technique for simultaneous multispectral imaging of combustion species. FRAME uses a non-conventional illumination scheme wherein each excitation source is given a unique periodic spatial code (line structure) that can be recognized and isolated by a spatial frequency-sensitive algorithm in the data post-processing. This methodology makes it possible for a single camera to collect several images at the same time. Past methods to achieve simultaneous multispectral imaging in combustion often rely on schemes based on sequential illumination and detection using several intensified, time-gated cameras to avoid signal interference between different signal components. By acquiring all data with a single detector using the FRAME concept, correction procedures to (1) achieve precise pixel-to-pixel overlap and (2) compensate for differences in the collection optics and detector characteristics are no longer needed. Laser sheet-based FRAME imaging does, however, require additional optical access due to the need to illuminate the sample at two angles. Moreover, the current approach to create the intensity modulation yields losses on the excitation side that needs to be addressed experimentally to improve the SNR. Lossless methods to create the periodic line structure using e.g. diffraction optical elements will be investigated in future work.

Since FRAME collects all data during a single acquisition and not in a rapid sequence, fast camera gating becomes less critical. This could potentially allow intensified cameras to be replaced with less expensive, yet more sensitive cameras that also have improved pixel resolution. The robustness of the methodology was investigated in terms of beam steering where no major complications were seen in the current experimental configuration. Furthermore, POD mode analysis was performed on FRAME data of OH and CH$_2$O with different image resolutions. The results indicate that the spatial resolution of the FRAME data that is achieved here is sufficient for POD mode analysis.

Finally, although FRAME was used here to differentiate between different fluorescence signals, the technique can readily be applied in a variety of alternative optical configurations, e.g. for measurements based on absorption, elastic scattering and polarization.

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References

[1] A.C. Eckbreth, Laser diagnostics for combustion temperature and species, Abacus, 1988.
[2] K. Kohse-Hoinghaus, R.S. Barlow, M. Alden, E. Wolfrum, Combustion at the focus: laser diagnostics and control, Proc. Combust. Inst. 30 (2005) 89–123.
[3] Y. Ishino, N. Ohiwa, Three-dimensional computerized tomographic reconstruction of instantaneous distribution of chemiluminescence of a turbulent premixed flame, JSME Int. J. B. Fluid. T. 48 (2005) 34–40.
[4] C. Gilbert, G. Lu, Y. Yan, Three-dimensional tomographic reconstruction of the luminosity distribution of a combustion flame, IEEE Trans. Instrum. Meas. 56 (2007) 1300–1306.
C. [58x595] 1165–1184

lumination

structures

A. [58x626] (2002)

J. [58x515] G. [58x618] C. [58x491] Combust. [58x666] Flame

B. [58x443] Flame

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S. [58x347] diluted

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R.J. [58x555] 159–169

Schulz

Roder

Adrian

Frank

Dyer

dilution

using laser diagnostic methods, Combust. Sci. Technol. 178 (2006)

1165–1184.

C. [58x595] 1165–1184

lumination

structures

A. [58x626] (2002)

J. [58x515] G. [58x618] C. [58x491] Combust. [58x666] Flame

B. [58x443] Flame

lease

S. [58x347] diluted

M. [58x332] CH-OHPLIF

R.J. [58x555] 159–169

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