



Article Similarity Analysis for Time Series-Based 2D Temperature Measurement of Engine Exhaust Gas in TDLAT

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Abstract: As regulations on the emission of pollutants from combustion systems are further tightened, it is necessary to reduce pollutant species and improve combustion efficiency to completely understand the process in the combustion field. Tunable diode laser absorption tomography (TDLAT) is a powerful tool that can analyze two-dimensional (2D) temperature and species concentration with fast-response and non-contact. In this study, stabilized spectra were implemented using the mean periodic signal technique to enable real-time 2D temperature measurement in harsh conditions. A time series statistical-based verification algorithm was introduced to select an optimal spectral cycle to track 2D reconstruction temperature. The statistical-based verification is based on the Two-sample *t* test, root mean square error, and time-based Mahalanobis distance, which is a technique for similarity analysis between thermocouple and reconstruction temperature of 18 candidate cycles. As a result, it was observed that the statistical-based TDLAT contribute to improving the accuracy of time series-based 2D temperature measurements.

Keywords: tunable diode laser absorption tomography; exhaust gas; temperature; similarity analysis; Mahalanobis distance; root mean square error

1. Introduction

With global warming prevention and environmental protection emerging as an issue all over the world, emission regulations are gradually being tightened as the effects of emissions from combustion systems on air and human bodies are highlighted. For example, EURO 6 regulations are an effort to reduce emissions such as CO, HC, NO_x, and PM by the auto industry. The recent EURO 6d emission regulations require real driving emissions (RDE) tests, which measure pollutants emitted from motor vehicles while they are driven, as an additional approval. As a result, demand for vehicles with low NO_x emissions and high engine power has increased, and several studies have been conducted to improve the performance of NO_x after treatment systems [1–3]. An important element in catalytic after treatment devices is exhaust gas temperature because catalytic performance is a function of temperature. A car engine running at high temperature tends to emit less CO₂ and more NO_x. On the other hand, a lower engine temperature will result in more CO_2 and less NO_x . However, the challenge of reducing both CO₂ and pollutant emissions simultaneously still remains. Therefore, analysis of temperature and species concentration during combustion is necessary for fundamental understanding of combustion reaction, which may reduce pollutants in the combustion system and improve combustion efficiency. However, the thermocouple (TC), a conventionally used one-point temperature measurement device, has difficulty in analyzing non-uniform flow, such as in engine exhaust. In addition, the techniques such as coherent anti-stokes Raman scattering (CARS) [4,5] and laser induced grating spectroscopy

(LIGS) [6], which are recently used in the spectroscopy field, can measure temperature at a fast response speed, but have a limit of point measurement, and filtered Rayleigh scattering (FRS) [7] and planer laser induced fluorescence (PLIF) [8,9] enable two-dimensional temperature, but their measurement systems are expensive and complicated. Tunable diode laser absorption tomography (TDLAT) is a suitable tool for understanding the combustion process by allowing 2D analysis of temperature and species concentration with non-contact fast-response. Therefore, it has been developed with various TDLAT techniques for measuring 2D temperature and species concentration in combustion fields [10–14]. Deguchi et al. demonstrated a 2D temperature distribution measurement by TDLAT on Bunsen-type flame burners and gasoline engines [10]. Ma et al. developed a hyperspectral tomography (HT) method that can simultaneously measure 2D distribution of temperature and H₂O concentration with a temporal resolution of 50 kHz at the exhaust plane of a j85 engine [11]. Busa et al. developed a measurement method of combustion efficiency that combines TDLAT and particle image velocimetry (PIV) in the scramjet engine [13]. However, these methods cannot be applied directly for real-time measurements because of excessive computational time and lack of time resolution assessment required for computed tomography (CT) analysis.

This study introduced a spectral cycle parameter improving reconstruction performance with a posteriori statistical technique in conventional absorption tomography to measure the exhaust gas temperature in milliseconds through TDLAT. Signal stabilization technique was applied in various cycle models to obtain stabilized absorption spectra in harsh environments, and statistical verification using TC measurement was performed to evaluate the results of the time series 2D reconstruction temperature distribution.

2. Theory of TDLAS

Tunable diode laser absorption spectroscopy (TDLAS) is a measurement technique commonly used in a variety of applications recently as well as temperature and density of molecules in gas [15,16]. When a laser beam permeates an absorption medium, its intensity will be attenuated by the absorbing gas species. This phenomenon is measured by TDLAS system that continuously scans laser wavelengths. The principle of TDLAS is theoretically expressed in terms of absorption by Beer-Lambert law that describes the relationship between the incident intensity I_{v0} and transmitted intensity I_v as follows:

$$\frac{I_v}{I_{v0}} = exp\{-A_v\} = exp\left\{-\int_0^L \alpha_v(s)ds\right\},\tag{1}$$

where A_{ν} denotes the spectral absorbance of representative wavenumber ν . *L* denotes the optical path length, and α_{ν} is the local mass-weighted absorption coefficient at position s along the laser beam, from 0 to *L*. This quantity is given by the following expression:

$$\alpha_{v} = \sum_{i} \frac{x_{i}p}{kT} \sum_{j} S_{i,j}(T) f_{v} \Big(v; v_{j}, \Delta v_{G}, \Delta v_{L} \Big),$$
⁽²⁾

where, k [J/K] denotes Boltzmann's constant, f_v is the Voigt profile, with Lorentzian and Gaussian half width at half maximums (HWHMs), Δv_L and Δv_G [17–19]. p denotes pressure, assumed to be uniform and x_i denotes mole-fraction of absorbing species i, where, x_i , p, and T are evaluated locally. $S_{i,j}(T)$ denotes temperature dependent line strength of absorption transition j and can be calculated by a function of temperature as follows:

$$S_{i,j}(T) = S_{i,j}(T_0) \frac{Q(T_0)}{Q(T)} \frac{T}{T_0} exp \left[-\frac{hc E_i''}{k} \left(\frac{1}{T} - \frac{1}{T_0} \right) \right] \left[1 - exp \left(\frac{-hc v_{0,i}}{kT} \right) \right] \left[1 - exp \left(\frac{-hc v_{0,i}}{kT_0} \right) \right]^{-1}, \quad (3)$$

where, h (J·s) and c (cm/s) denote Planck's constant and the speed of light, v_0 (cm⁻¹) denotes the line-center wavenumber, E_i'' (cm⁻¹) denotes lower-state energy of the transition, T_0 (K) is the reference

temperature (296 K), and Q(T) denotes the partition function of molecular absorption at a particular temperature. It is usually expressed as a 4th order polynomial function [20]. Although it is difficult to accurately describe the chemical formula of gasoline, iso octane (C₈H₁₈) was selected because it is the hydrocarbon component with similar thermal properties to the actual composition. The combustion reaction of iso octane is as follows:

$$C_8H_{18} + 12.05(O_2 + 3.76N_2) \rightarrow 8CO_2 + 9H_2O + 12.05 \cdot (3.76)N_2.$$
 (4)

Among the combustion products, nitrogen (N_2) is an inert gas and hence, there is no chemical or absorption reaction. Therefore, water-vapor (H_2O) was chosen for exhaust gas diagnosis because of its high mole fraction and high absorption strength.

Figure 1 shows the theoretical H₂O absorption spectra calculated using the HITRAN2012 molecular spectroscopic database [21]. The patterns of the absorption spectra can be observed as the temperature changes. In this study, three peaks lines (7202.255 cm⁻¹, 7202.909 cm⁻¹, and 7203.891 cm⁻¹) with high line strengths and no interference with other molecules were chosen for calculating reconstruction temperature.



Figure 1. Theoretical H₂O absorption spectrum.

3. Experiment and Tomography Analysis

A schematic diagram of a TDLAS system measuring engine exhaust temperature is shown in Figure 2. The OHC (over head camshaft) gasoline engine (EX17, Subaru Robin, Doylestown, PA, USA) is driven in idle mode to produce emissions. The exhaust pipe installed 5 mm below the measurement cell was 20 mm in diameter and 150 mm in length. The distributed feedback (DFB) laser (NLK1E5GAAA, NTT Electronics, Japan) for H₂O gas target is tuned to the near-infrared spectral region at 1388 nm by a laser diode controller (LDC-3900, ILX Lightwave, Bozeman, MT, USA). The modulated laser beam is split into 16 channels by a 1 × 16 fiber optic coupler (SMF-28e, OPNETI, NY, USA). Among them, the 10 beams are irradiated into 10 collimators (50-1310-APC, THORLABS, Newton, NJ, USA) and the transmitted optical signals are detected by 10 photodetectors (G12180-010A, Hamamatsu Photonics, Japan). The received laser signals are amplified and filtered by low pass filters to remove signal noise. The amplified signals are transmitted into a data recorder (8861 Memory High Coda HD Analog16, HIOKI, Japan) in real time at a sampling rate of 500 kHz. Simultaneously, the real temperature for verification is measured by a B-type TC at the sampling rate of 1 kHz at the center of the target area. After absorption, signals collect by the data acquisition card are sent to the computer for reconstruction calculation.

Multiple parallel laser beams pass through the region of interest. The intersected spectral information is used to reconstruct 2D temperature by CT analysis. Figure 3 shows the laser paths and grids in the measurement region. The laser path consists of 10 real beams in a 5×5 layout and eight interpolated virtual beams The interpolated virtual beams imply the virtual signal data interpolated from the signal of the actual beam, and in the signal application field a widely introduced cubic spline interpolation is used in this study [22]. The cell length through which the laser passes in the measurement area is 60 mm and reconstruction area is 40 mm, and the spatial region for CT analysis consist of 81 (9 × 9) square grids with a grid size of 2.0 mm × 2.0 mm.



Figure 2. Schematic diagram of the experimental setup using tunable diode laser absorption spectroscopy (TDLAS).



Figure 3. Laser paths and measurement cell.

Unexpected error signals due to harsh conditions were removed to reduce reconstruction calculation time for converging. The formula is as follows:

$$\delta_t = \sum_{n=1}^N \frac{\left| \bar{I}_{v0,i} - \bar{I}_{v,i} \right|_t}{\left| \bar{I}_{v0,i} - \bar{I}_{v,i} \right|_{t-1}},\tag{5}$$

where, the subscript *t* denotes a cycle at *t* time and *n* denotes the data number in a cycle. δ denotes the error rate, which is a ratio of absolute error sum in \bar{I}_v and \bar{I}_{v0} of the cycle to that of the previous cycle. Here, the signal cycle was eliminated when δ_t is greater than 1.5. This method effectively reduced the error when converting the signal into the spectra. The noises generated by beam steering, window foul, and thermoelectric noises were approximated by Gaussian noise modeling, and the stabilized absorption spectra was obtained by the polynomial reduction method (PRM) averaging a number of absorption signal cycles [10,23,24] Although averaging a significant number of absorption cycles provides more stable signal, it has the drawback of degrading time-resolution and distorting raw signals. Hence, it is necessary to investigate appropriate number of cycles with good reconstruction performance and fast time-resolution simultaneously.

Figure 4 shows the result of mean transmitted signals (\bar{I}_v) and mean absorption spectra using a total of 201 cycles (1 reference cycle, 100 cycles in front and 100 cycle in back) measured by the TDLAS system. Here, one cycle of measured signal consists of 500 samples with a recording time of 1 ms. A total of 5 million signals were received during 10 s of engine operation. Figure 4a is the result of averaging each raw signal acquired from the photodetector at the same time when 10 lasers passed through the target area. To convert these signals into absorption spectra, \bar{I}_v and the mean incident signal (\bar{I}_{v0}) must be known as shown in Equation (4). \bar{I}_{v0} , also known as the baseline, was estimated by PRM, which is a least squares fitting using 5th order polynomial curve with \bar{I}_v [23,25]. To facilitate curve fitting, the saw-tooth waveform had a frequency of 1 kHz and a sampling rate of 2 µs. All measured absorption spectra generated from \bar{I}_v and \bar{I}_{v0} is shown in Figure 4b. These spectra consisted of 311 wavenumbers. Laser group 0 and 1 represent a group of vertical and horizontal lasers, respectively. The spectra pattern can be identified by different laser numbers. As shown in Figure 1, the measured absorption spectra are determined by the temperature of the measurement area, so the reconstructed temperature distribution in measurement zone can be traced by a tomographic reconstruction method from the information of each absorption spectrum.



Figure 4. Results of mean transmission signals and mean absorption spectra of 10 laser beams: (a) 10 mean transmitted signals using 201 cycles and (b) 10 mean absorption spectra using 201 cycles.

The determination method of cycle and temperature is illustrated in Figure 5. First, raw absorption signals, reference time *i* and candidate cycle *j* are input. *i* denotes analysis time point at engine running time. A total of 990 absorption signals were used in 0.1 s from 1.0 to 9.9 s. *j* denotes the number of cycles of absorption signal to be averaged. Eighteen candidate cycles were chosen to evaluate the analytical performance in various cycles (31, 51, 71, 91, 111, 151, 201, 251, 301, 351, 401, 451, 501, 601, 701, 801, 901, and 1001 cycles).



Figure 5. Flowchart for tracking cycle and reconstructed temperature using TDLAT.

In the second step, the PRM and δ value are combined to obtain mean absorption spectra, described in more detail above. This technique was able to remove noise more effectively than the conventional PRM. Reconstruction calculation consisting of 9 × 9 array was calculated using mean absorption spectra. The reconstructed temperature distribution was tracked by repeated calculations until errors in the experimental and theoretical absorption spectra were minimized by the multiplicative algebraic reconstruction technique (MART) algorithm [24].

$$\alpha_{v(n,m)}^{(k)} = \alpha_{v(n,m)}^{(k-1)} \times \left[\frac{A_v}{L_{(n,m)} \sum_{m=1}^M \sum_{n=1}^N \alpha_{v(n,m)}^{(k-1)}} \right]^{\beta L_{(n,m)}},$$
(6)

where, the superscript *k* represents the number of iteration and the subscript (n, m) is the gird number. β is the relaxation parameter that affects the rate of convergence and was set to 0.1. The initial values of $\alpha_{v(n,m)}$ at each grid are an important factor for fast convergence calculation, which were approximated to target values by adopting the multiplication line of sight (MLOS) method [26]. The reconstruction calculation was terminated when difference of $\alpha_{v(n,m)}^{(k)}$ and $\alpha_{v(n,m)}^{(k-1)}$ was less than 1×10^{-6} .

Figure 6 shows a comparison of the measured absorption spectra with the calculated absorption spectra in the engine running time of 5.0 s. The average root-mean-square error (RMSE) at 311 wavenumber were 9.766×10^{-4} , 4.485×10^{-5} , 1.273×10^{-3} , and 2.144×10^{-3} , respectively, in Figure 6a–d. Measured spectra were in good agreement with spectra by the theoretical calculation, but the measured spectra for the 51 and 201 cycles were slightly different from each other, and their reconstruction calculations were also differed slightly. This implies that the number of mean absorption cycles is an important parameter for the reconstruction results.

In the third step, the reconstructed temperatures, $T_{Rec(i,j)}$, calculated for each candidate cycle and time are output. To estimate the $T_{Rec(i,j)}$ from the finally converged $\alpha_{v(n,m)}$ at all grids, the theoretical absorption coefficients at the temperature has already been calculated using the HITRAN database for the mole fraction 1.0. The correlation between the theoretical absorption coefficient, $\alpha_{v,the}$, and the reconstructed absorption coefficient, α_v , was evaluated to find the value with the most similar pattern, i.e., the nearest correlation coefficient to 1.0, and to obtain $T_{Rec(i,j)}$. The correlation coefficient, $C_{n,m}$, at each grid is described as follows.

$$C_{n,m} = \frac{\sum_{v=1}^{n} (\alpha_v - \overline{\alpha}) (\alpha_{v,the} - \overline{\alpha}_{the})}{\sqrt{\sum_{v=1}^{n} (\alpha_v - \overline{\alpha})^2 \sum_{v=1}^{n} (\alpha_{v,the} - \overline{\alpha}_{the})^2}},$$
(7)

where $\overline{\alpha}$ and $\overline{\alpha}_{the}$ denote the average value each of the theoretical and reconstructed absorption coefficient.

Figure 7 shows the results of 2D reconstructed temperature distribution using 51 and 201 cycles at the engine running time of 5.0 s. Comparing Figure 7a,b, the temperature difference at the center point was 12.35 K and the RMSE at all grids was 5.87 K. This is an example of the highest temperature difference compared to other engine running times. The characteristics of the temperature distribution for each time were similar, but the temperature fluctuation was shown differently. From these results, it was found that the number of mean cycles affected the results of CT analysis. Additionally, the output \overline{T}_{Rec} was set to the average value of nine grids located in the center of the measurement area as the TC was installed in the center of the exhaust pipe and the temperature was measured at one point. After all, the output T_{Rec} consists of a set of elements \overline{T}_{Rec} for each i and j. The number of T_{Rec} was 17820 in the i × j array (990 × 18).

0.1

0.09

0.08

0.07

0.06

0.05

0.04

0.03

0.02

0.01

0.11

0.1

0.09

0.08

0.07

Absorption (a.u.)

0

7201.5

Absorption (a.u.)





(c)

(**d**)

Figure 6. The spectra of reconstructed and measured absorption data at the engine running time 5.0 s: (**a**) 51 cycles, boundary; (**b**) 201 cycles, boundary; (**c**) 51 cycles, center; and (**d**) 201 cycles, center.



Figure 7. 2D reconstructed temperature distributions at the engine running time 5.0 s: (**a**) 51 cycles and (**b**) 201 cycles.

In the fourth step, the statistical-based analysis was performed using T_{Rec} and T_{Thc} , which means temperature measured by TC and was used to determine the cycle and T_{Rec} most similar to T_{Thc} . Further details are described in Section 4.

The computation time of each step on an Intel Core i7 2600K CPU with 16GB RAM was as follows. The CPU time of Process I to convert signal data into mean absorption spectra was 1.38 s and 1.90 s (311 wavenumbers) respectively at 51 and 201 cycles. As the number of mean cycles increased, the CPU time increased linearly. The CPU time of Process II was scored in under 1.79 s (311 wavenumbers), mostly converging on less than 150 iterations.

4. Statistical-Based Mean Difference Test and Similarity Analysis

The statistical-based analysis is performed to identify cycles that represent T_{Rec} similar to T_{Thc} . The 18 candidate cycles were selected using suitable intervals for analysis from 31 to 1001 cycles. The procedure is shown in Figure 8. Paired *t* test, which is a mean difference testing method, is conducted to evaluate the similarity between the 18 candidate cycles [27].



Figure 8. Procedure for mean different test and similarity analysis.

The next step is followed by performing a mean difference test and similarity analysis between T_{Thc} and T_{Rec} for all 18 candidate cycles. The mean difference test uses the representative statistical technique of Two-sample *t* test. RMSE and time-based Mahalanobis distance (TMD) were used for similarity analysis. RMSE is a measure of generalization of standard deviations that indicates the difference between actual and estimated values [28]. The Mahalanobis distance (MD) represents the distance between data and considers covariance [29].

Finally, the cycles that represent T_{Rec} most similar to T_{Thc} based on the analysis results were determined.

4.1. [Step 1] Mean Difference Test

 T_{Rec} derived from 18 candidate cycles consists of 0.1 s intervals from 1 to 9.9 s as shown in Table 1. To test the mean temperature difference in 18 candidate cycles, the analysis was performed on nine $T_{Rec(i,j)}$, which is the reconstructed temperature of *i*th reference time of the *j*-cycles. The total number of sets comparing $T_{Rec(i,j)}$ in 18 candidate cycles to each other for the paired *t* test were 153. For the actual test, the analysis was performed only at the reference time (1 s, 2 s, 3 s, 4 s, 5 s, 6 s, 7 s, 8 s, and 9 s) for all 153 sets.

4.1.1. Paired *t* Test

The paired *t*-test is useful for analyzing the same set of items that were measured under two different conditions, i.e., differences in measurements made on the same subject before and after a treatment, or differences between two treatments given to the same subject.

The paired *t*-test is based on the test statistic as shown below:

$$T_{pt} = \frac{\overline{D} - d_0}{S_D / \sqrt{n}},\tag{8}$$

where,

$$\overline{D} = \frac{\sum_{k=1}^{n} \left(T_{Rec(i,A)^k} - T_{Rec(i,B)^k} \right)}{n},\tag{9}$$

$$S_D = \sqrt{\frac{\sum_{k=1}^{n} \left(T_{Rec(i,A)^k} - T_{Rec(i,B)^k} - \overline{D} \right)^2}{n-1}},$$
(10)

where, \overline{D} denotes the mean of the differences between the two kinds of cycles. d_0 denotes a constant that can be set depending on whether the mean of the difference between the two groups is zero or a specific non-zero value. S_D denotes the standard deviation of differences between two kinds of cycles. $T_{Rec(i,A)}^k$ denotes the *k*th reconstruction temperature at *i*th time (s) of cycle A, $i = 1.0, 1.1, 1.2, \cdots, 9.9$. $T_{Rec(i,B)}^k$ denotes the *k*th reconstruction temperature at *i*th time (s) of cycle B. *n* denotes the number of T_{Rec} in *i*th time (s).

Table 1. T_{Rec} per 18 candidate cycles used in [Step 1].

No.	Time			T _{Rec}		
1100	(s)	31	51	•••	901	1001
1		404.833	400.000		405.432	404.189
2		406.775	401.665		407.081	405.842
3		404.947	399.784		405.436	404.455
4		407.458	401.846		407.028	405.633
5	1.0	409.400	403.506		408.581	407.223
6		407.556	401.614		406.820	405.720
7		404.951	399.982		405.459	404.137
8		406.842	401.604		406.900	405.606
9		404.999	399.722		405.051	405.159
	•••			•••		•••
806		398.712	399.182		400.164	400.543
807		397.870	397.965		402.270	402.782
808	9.9	396.147	396.544		401.184	401.859
809		397.183	397.668		401.351	401.646
810		396.221	396.327		403.357	403.785

For paired *t* test, the hypothesis is as follows:

- Null hypothesis: H_0 : $\overline{D} = d_0$ (the population mean of the difference (\overline{D}) equals the hypothesized mean of the difference ($d_0 = 0$).
- Alternative hypothesis: H_1 : $\overline{D} \neq d_0$ (the population mean of the difference (\overline{D}) does not equal the hypothesized mean of the difference ($d_0 = 0$).

Here, the null hypothesis is rejected when the *p*-value is less than or equal to 0.05, i.e., when there is no difference between the mean of the two groups.

4.1.2. Results of Mean Difference Test in [Step 1]

Table 2 is a summary of the paired *t*-test results. For 153 sets, 1377 test results would be obtained when each test is performed from 1 to 9 s. If the *p*-value is less than 0.05, the null hypothesis is rejected. The *p*-values of 1 s, 3 s, 8 s, and 9 s were less than 0.05, i.e., the means of all cases were not equal to each other. In 2 s, the *p*-value was less than 0.05 except in four sets (i.e., 251 vs. 301, 451 vs. 501, 451 vs. 601, and 501 vs. 601). In 4 s, only one set (351 vs. 401) showed a *p*-value greater than 0.05 and the remaining 152 sets had a *p*-value of less than 0.05. In 5 s, 6 s, and 7 s, the *p*-value was greater than 0.05, while the remaining 152 sets had a *p*-value less than 0.05. Therefore, except for eight sets among 1377 sets, the *p*-value in all sets was less than 0.05 and hence, there was a mean difference between the cycles.

Time (s)		Test Sets		<i>p</i> -Value
1.0		All sets		< 0.05
	251	vs.	301	0.626
2.0	451	vs.	501	0.476
2.0	451	vs.	601	0.402
	501	vs.	601	0.331
3.0		All sets		< 0.05
4.0	351	vs.	401	0.371
5.0	451	vs.	501	0.208
6.0	401	vs.	451	0.633
7.0	601	vs.	701	0.221
8.0		All sets		< 0.05
9.0		All sets		< 0.05

Table 2. Results of paired *t*-test.

4.2. [Step 2] Similarity Analysis

Table 3 shows the $T_{Thc(i)}$ and $\overline{T}_{Rec(i,j)}$ of all 18 candidate cycles used in the analysis where each $\overline{T}_{Rec(i,j)}$ is the mean value for nine $T_{Rec(i,j)}^{k}$ per 0.1 s interval in Table 1. For example, in Table 3, $\overline{T}_{Rec(1.0,31)}^{9}$ = 406.418, which averaged reconstruction temperatures from $T_{Rec(1.0,31)}^{1}$ = 404.833 to $T_{Rec(1.0,31)}^{9}$ = 404.999 in Table 1. Here, three methods were used for similarity analysis. First, the Two-sample *t* test was used to test the difference between the means of the two groups. Second, RMSE analysis was performed to address the difference between estimates or models and observed values in real-world settings. RMSE analysis can be used to aggregate the differences between the residuals of two groups into one measure. Finally, in TMD analysis the correlation between two groups with different means was taken into account.

		Time	T	$\overline{\overline{T}}$					
No.	Group	(s)	T _{Thc(i)}	31	51		1001		
1		1.0	408.000	406.418	401.080		405.204		
2		1.1	408.688	402.722	401.324		406.271		
3		1.2	409.921	407.890	407.890		406.525		
4		1.3	407.798	410.285	411.842		406.990		
5	1	1.4	411.453	402.434	411.723		406.453		
6	1	1.5	406.759	392.685	401.679		406.284		
7		1.6	406.005	397.013	405.508		407.419		
8		1.7	405.461	418.457	416.981		407.537		
9		1.8	405.110	407.200	401.237		406.350		
10		1.9	404.842	416.607	407.599		405.416		
80		9.0	404.658	397.217	397.462		401.795		
81		9.1	405.853	394.707	393.932		401.638		
82		9.2	402.657	408.041	403.190		400.741		
83		9.3	400.688	395.484	396.351		400.231		
84	0	9.4	401.921	402.101	398.922		401.280		
85	9	9.5	399.798	393.647	396.263		401.277		
86		9.6	403.453	410.205	399.067		401.559		
87		9.7	400.895	390.764	398.329		401.343		
89		9.8	403.987	398.441	401.295		400.894		
90		9.9	401.752	403.739	409.931		400.857		

Table 3. Temperatures used in [Step 2].

4.2.1. Two-Sample *t*-Test

Two-sample *t*-test is a widely used hypothesis test used to compare whether the averages are equal. The assumptions made while doing a Two-sample *t*-test include: (i) the data are continuous, (ii) the data follows normal probability distribution, (iii) the variances of the two populations are equal, (iv) two-sample is independent, and (v) both samples are simple random samples from their respective populations. The Two-sample *t*-test is based on the test statistic shown below:

$$T_{tw} = \frac{\overline{T}_{Thc^{g}} - \overline{\overline{T}}_{Rec^{g}}}{\sqrt{T_{S^{2}}}} \sqrt{\frac{n_{T^{g}} n_{R^{g}}}{n_{T^{g}} + n_{R^{g}}}},$$
(11)

$$T_{S^2} = \frac{\sum_{i=1}^{n_{T^S}} \left(T_{Thc(i)} - \overline{T}_{Thc^S} \right)^2 + \sum_{i=1}^{n_{R^S}} \left(\overline{T}_{Rec(i,j)} - \overline{T}_{Rec^S} \right)^2}{n_{T^S} + n_{R^S} - 2},$$
(12)

where, T_{S^2} denotes the variance of $T_{Thc(i)}$ and $T_{Rec(i,j)}$. $T_{Thc(i)}$ denotes the TC temperature at *i*th time (s), $i = 1.0, 1.1, 1.2, \dots, 9.9$. $\overline{T}_{Thc}{}^{g}$ denotes the mean of all $T_{Thc(i)}$ in the *g*th group, $g = 1, 2, \dots, 9$. $\overline{\overline{T}}_{Rec}{}^{g}$ denotes the mean of all $\overline{T}_{Rec(i,j)}$ in the *g*th group. n_{Tg} denotes the number of all $T_{Thc(i)}$ in the *g*th group. $n_{Rec}{}^{g}$ denotes the number of all $T_{Rec(i,j)}$ in the *g*th group.

For the Two-sample *t* test, the hypothesis is as follows

- Null hypothesis: $H_0: = \overline{T}_{Thc}{}^g \overline{\overline{T}}_{Rec}{}^g = \delta_0$ (the difference between the population means $(\overline{T}_{Thc}{}^g \overline{\overline{T}}_{Rec}{}^g)$ is equal to the hypothesized difference $(\delta_0 = 0)$).
- Alternative hypothesis: $H_1: = \overline{T}_{Thc}{}^g \overline{\overline{T}}_{Rec}{}^g \neq \delta_0$ (the difference between the population means $(\overline{T}_{Thc}{}^g \overline{\overline{T}}_{Rec}{}^g)$ is not equal to the hypothesized difference $(\delta_0 \neq 0)$).

Here, the null hypothesis is rejected when the p-value is less than or equal to 0.05, i.e., when there is no difference between the mean of the two groups. In this study, statistical software version 18 of Minitab was used for the Two-sample t test.

4.2.2. Root Mean Square Error (RMSE)

RMSE is used to measure the difference between values predicted by a model or an estimator and the value observed. The RMSE is a measure of accuracy to compare forecasting errors of different models for a particular dataset and not between the dataset as it is scale-dependent. RMSE is always non-negative and a value of zero would indicate a perfect fit to the data. In general, a lower RMSE is better than a higher one. The formula is as follows:

$$T_{RMSE} = \sqrt{\frac{1}{m} \times \sum_{i=1}^{m} (T_{Thc(i)} - T_{Rec(i,j)})^2}.$$
 (13)

4.2.3. Time-Based Mahalanobis Distance (TMD)

MD was introduced by P. C. Mahalanobis in 1936 to measure distance between a point P and distribution D. It is a multi-dimensional generalization of the idea for measuring the amount of standard deviations P is away from the mean of D [30]. Among various methods for measuring distance (i.e., Euclidean distance, Manhattan distance, etc.), MD is used as it has the advantage of calculating the distance by considering weight according to the magnitude of covariance between variables [31]. Thus, MD is a method that is used to assess similarities between two groups. In this study, TMD analysis is performed to analyze similarities between T_{Thc} and \overline{T}_{Rec} by reference time. The closer the TMD value is to zero, the more likely T_{Thc} and \overline{T}_{Rec} are to be similar. The calculation of TMD procedure is as follows:

[Stage 1] Calculate the covariance matrix (S) and TMD using the following formula (14) and (15).

$$S = \begin{bmatrix} Var[T_{Thc(i)}] & Cov[T_{Thc(i)}, \overline{T}_{Rec(i,j)}] \\ Cov[\overline{T}_{Rec(i,j)}, T_{Thc(i)}] & Var[\overline{T}_{Rec(i,j)}] \end{bmatrix},$$
(14)

$$MD = \sqrt{\left(X_i - \overline{X}\right)^T S^{-1} \left(X_i - \overline{X}\right)},\tag{15}$$

$$(X_i - \overline{X}) = \begin{pmatrix} T_{Thc(i)} - \overline{T}_{Thc} \\ = \\ \overline{T}_{Rec(i,j)} - \overline{T}_{Rec(i,j)} \end{pmatrix},$$
 (16)

where, \overline{T}_{Thc} denotes the geometric mean of all $T_{Thc(i)}$, $i = 1.0, 1.1, 1.2, \dots, 9.9$. $\overline{\overline{T}}_{Rec(i,j)}$ denotes the geometric mean of all $\overline{T}_{Rec(i,j)}$ in *j*th cycles. S⁻¹ denotes inverse covariance matrix. *T* indicates that the vector should be transposed.

[Stage 2] The obtained TMD values are calculated using the following formula for each time (s) interval.

$$T_{MD} = (MD_1 \times MD_2 \times \dots \times MD_n)^{\frac{1}{n}}, \ n = 1, 2, \cdots m.$$
(17)

4.2.4. Results of Similarity Analysis in [Step 2]

Two-sample *t* test, RMSE, and TMD analysis were performed to select the cycles that indicate a T_{Rec} similar to that of T_{Thc} . The results of a normality test conducted on T_{Thc} and T_{Rec} for 18 candidate cycles prior to the Two-sample *t* test showed that all were satisfactory for normality. However, the equivalence test between T_{Thc} and T_{Rec} showed that some results did not satisfy the equivalence. The results of the equivalence test are summarized in Appendix A. In general, Welch's *t*-test is used if the two groups do not meet equal variances. However, if the very small samples (n < 10), the Two-sample *t* test performs slightly better than Welch's *t*-test [32].

The first Two-sample *t*-test results are shown in Table 4 and the null hypothesis is rejected when the *p*-value is less than 0.05 (see Section 4.2.1). The analysis results are explained based on the cycle, all *p*-values from 1 to 9 s for 31, 51, 91, 111, and 251 cycles were above 0.05. For the 71 cycles, the *p*-value was close to 1 at 2 s, 5 s, 6 s, and 8 s. In other words, the probability of selecting null hypothesis was high. For 151 and 201 cycles, the *p*-value was over 0.05 for all seconds except 5 s. On the other hand, the 451 cycles showed *p*-value less than 0.05 at 5 s and 9 s. The *p*-value of the 901 cycles was less than 0.05 at 8 s and 9 s. In the 501, 601, 701, and 801 cycles, the *p*-values were less than 0.05 for 5 s, 8 s, and 9 s. In the case of the 1001 cycles, the *p*-value of 3 s, 8 s, and 9 s was found to be below 0.05.

The result of RMSE analysis is shown is Table 5. The smaller the RMSE values, the more the similarity between T_{Thc} and T_{Rec} . First, the analysis results were based on 10 temperatures per group from 1 to 9 s (see Table 3). In terms of time, the RMSE value (1.951) of 401 cycles in 1 s was the smallest. In 2 s, the RMSE value of 1001 cycles was the smallest at 2.270. RMSE values for 601 cycles were the smallest in 3 s and 4 s. The RMSE values for 251, 201, and 701 cycles were the lowest in 5 s, 6 s, and 7 s. RMSE values for 1001 cycles were the smallest in 8 s and 9 s. The analysis using 90 temperatures from 1 to 9.9 s (see Table 3) showed that the RMSE value of 601 cycles was the smallest.

Lastly, the analysis results of TMD are shown is Table 6. The smaller the TMD value, the more the similarity between T_{Thc} and T_{Rec} . The analysis results were based on 10 temperatures per group from 1 to 9 s (see Table 3). In terms of time, the TMD value (1.626) of 351 cycles in 1 s was the smallest. In 2 s, the TMD value of 111 cycles was the smallest at 0.770. The TMD value for 151, 201, 251, and 801 cycles were the lowest in 3 s, 4 s, 5 s, and 6 s. TMD values for 201 cycles were the smallest in 7 s, 8 s, and 9 s. Next, the analysis using 90 temperature readings from 1 to 9.9 s showed that the TMD value (0.933) of 201 cycles was the smallest.

	Time (s)												
Cycles	1	2	3	4	5	6	7	8	9				
-					<i>p</i> -Value								
31	0.647	0.463	0.839	0.273	0.692	0.496	0.924	0.976	0.169				
51	0.711	0.726	0.316	0.292	0.707	0.681	0.864	0.728	0.069				
71	0.774	0.985	0.147	0.292	0.982	0.896	0.727	0.997	0.164				
91	0.603	0.891	0.225	0.441	0.369	0.985	0.842	0.693	0.362				
111	0.531	0.983	0.390	0.667	0.092	0.901	0.828	0.352	0.482				
151	0.486	0.892	0.402	0.671	< 0.05	0.779	0.708	0.574	0.474				
201	0.396	0.964	0.328	0.381	< 0.05	0.958	0.684	0.488	0.177				
251	0.510	0.858	0.271	0.289	0.074	0.743	0.791	0.467	0.078				
301	0.384	0.890	0.246	0.325	< 0.05	0.789	0.928	0.137	0.109				
351	0.415	0.928	0.196	0.411	< 0.05	0.770	0.979	0.102	0.073				
401	0.406	0.790	0.177	0.253	< 0.05	0.655	0.984	0.112	0.051				
451	0.530	0.735	0.136	0.235	< 0.05	0.537	0.946	0.081	< 0.05				
501	0.509	0.691	0.136	0.269	< 0.05	0.622	0.865	< 0.05	< 0.05				
601	0.478	0.620	0.097	0.284	< 0.05	0.495	0.886	< 0.05	< 0.05				
701	0.432	0.631	0.084	0.367	< 0.05	0.449	0.911	< 0.05	0.054				
801	0.357	0.550	0.058	0.336	< 0.05	0.430	0.977	< 0.05	< 0.05				
901	0.316	0.574	0.053	0.431	0.062	0.431	0.871	< 0.05	< 0.05				
1001	0.216	0.528	< 0.05	0.446	0.057	0.413	0.773	< 0.05	< 0.05				

Table 4. Results of Two-sample *t*-test.

Table 5. Results of root mean square error (RMSE) analysis between T_{Thc} and T_{Rec} for 18 candidate cycles.

		Total									
Cycles	1	2	3	4	5	5 6 7		8	9	- 10001	
					RMSE					RMSE	Rank
31	8.487	8.039	10.325	8.441	12.622	7.869	7.320	5.887	5.887	8.578	18
51	5.514	5.592	6.370	5.994	7.759	4.776	6.912	5.526	5.526	6.049	17
71	6.269	4.497	5.404	6.334	5.316	6.007	6.028	4.420	4.420	5.538	16
91	4.879	3.775	4.331	5.559	3.894	5.921	5.073	3.620	3.620	4.822	15
111	4.702	3.860	4.282	5.046	3.580	5.265	4.156	3.496	3.496	4.393	14
151	3.755	3.129	4.216	4.833	3.564	2.957	4.314	4.065	4.065	3.918	13
201	3.177	2.978	3.349	3.376	2.900	2.529	3.810	3.736	3.736	3.199	12
251	3.024	3.063	3.496	3.400	2.338	3.166	3.243	2.755	2.755	3.120	11
301	2.603	2.884	3.159	2.961	2.749	2.844	2.541	2.267	2.267	2.809	10
351	2.056	2.733	2.799	2.686	2.955	3.333	2.529	2.099	2.099	2.703	9
401	1.951	2.583	2.641	2.451	2.849	3.149	2.546	2.456	2.456	2.584	4
451	2.047	2.909	2.662	2.403	2.785	3.439	2.512	2.156	2.156	2.641	7
501	2.433	2.580	2.703	2.278	2.894	3.543	2.433	2.140	2.140	2.643	8
601	2.096	2.453	2.561	2.265	2.947	3.368	2.182	2.039	2.039	2.491	1
701	2.201	2.591	2.899	2.426	2.925	3.544	1.905	1.987	1.987	2.555	3
801	2.209	2.318	3.055	2.629	2.771	3.492	1.907	1.870	1.870	2.549	2
901	2.438	2.499	3.174	2.568	2.735	3.693	2.054	1.715	1.715	2.624	6
1001	2.434	2.270	3.438	2.576	2.693	3.606	2.041	1.660	1.660	2.597	5

		То	tal								
Cycles	1	2	3	4	5	6	7	8	9	- 10	tui
-					TMD					TMD	Rank
31	1.860	1.124	1.188	1.208	1.174	1.383	0.889	0.731	0.880	1.121	4
51	1.758	1.079	1.279	1.267	1.204	1.253	0.910	0.854	0.876	1.136	6
71	1.829	0.967	1.250	1.305	0.995	1.363	0.970	0.736	1.014	1.123	5
91	1.780	0.873	1.170	1.348	1.006	1.299	0.915	0.773	1.148	1.112	3
111	1.777	0.770	1.155	1.307	1.022	1.304	0.921	0.835	0.962	1.082	2
151	1.727	0.936	1.133	1.456	1.143	1.303	1.052	0.997	1.071	1.181	10
201	2.132	0.798	1.552	0.888	1.549	0.992	0.684	0.429	0.506	0.933	1
251	1.675	1.277	1.391	1.273	0.965	1.361	1.016	0.812	1.163	1.190	14
301	1.719	1.271	1.506	1.252	1.258	1.291	1.050	0.755	0.872	1.185	12
351	1.626	1.280	1.394	1.261	1.291	1.481	0.996	0.745	0.911	1.188	13
401	1.708	1.297	1.477	1.273	1.363	1.263	1.046	0.911	0.993	1.237	18
451	1.772	1.375	1.312	1.292	1.205	1.258	0.972	0.788	1.033	1.194	16
501	1.871	1.342	1.264	1.328	1.309	1.153	0.945	0.811	1.056	1.200	17
601	1.880	1.373	1.262	1.439	1.320	0.981	0.997	0.726	1.098	1.191	15
701	1.956	1.355	1.341	1.458	1.258	0.990	0.882	0.754	1.053	1.182	11
801	2.012	1.320	1.347	1.479	1.299	0.849	0.814	0.727	1.147	1.165	8
901	2.061	1.319	1.411	1.400	1.246	1.055	0.779	0.693	1.138	1.177	9
1001	2.058	1.233	1.382	1.393	1.243	1.006	0.788	0.655	1.153	1.154	7

Table 6. Results of time-based Mahalanobis distance (TMD) analysis between T_{Thc} and T_{Rec} for 18 candidate cycles.

5. Conclusions

The results of this study are summarized as follows: (*i*) the paired *t* test for 18 candidate cycles indicated that, in most cases, T_{Rec} was not the same in each cycle. This implies that T_{Rec} might vary depending on the cycle set during CT analysis. Therefore, a standard for proper cycle setting is needed. (*ii*) Three types of similarity analysis yielded different results. First, the Two-sample t test showed that the T_{Rec} for 71 cycles was not the much different from that of T_{Thc} . Second, the results from RMSE showed that the T_{Rec} of the 601 cycles was most similar to that of the T_{Thc} . Finally, TMD results show that the T_{Rec} for 201 cycles was most similar to that of T_{Thc} . (iii) The reason for different results is that all three methodologies have different characteristics. For a more visualized description, a comparison of T_{Thc} and T_{Rec} was provided based on the similarity analysis result. Figures 9–11 show a comparison of the T_{Thc} with the T_{Rec} for each 71, 601, and 201 cycles, respectively. The movement of T_{Thc} and T_{Rec} over time shows that T_{Rec} for 601 and 201 cycles moved closely with T_{Thc} . However, a closer look at the temperature change pattern shows that T_{Rec} of 201 cycles was moving similarly to T_{Thc} . On the other hand, for 71 cycle, the range of T_{Rec} changes over time was observed to be much larger than T_{Thc} . (iv) For the Two-sample t-test, it was difficult to determine similarity by sensitively reflecting changes in temperature as the difference in the mean of the overall temperature data in a given period was taken into account. Likewise, the RMSE was purely an indicator of the mean of distance between T_{Thc} and T_{Rec} , which made it difficult to analyze, including the correlation between changes in T_{Thc} and T_{Rec} . TMD was a suitable indicator for this study because it considers both the distance and correlation between T_{Thc} and T_{Rec} . Therefore, if the engine used in this study is in idle mode, setting it to 201 cycles in the CT analysis will increase the accuracy of the analysis while minimizing the analysis time and effort.

Figure 12 shows the 2D reconstruction temperature distribution in 201 cycles in time from 1.0 to 9.0 s. The temperature of the peak area of the 2D distribution in each time was equal to that of the corresponding time at the 201 cycle reconstruction temperature in Figure 11. As illustrated in Figure 12, it can be seen that the maximum temperature was continuously distributed in the central area. For example, Figure 12b,h show a reconstruction temperature of approximately 410 K and 408 K

in the central area, and Figure 10 shows also the same temperature at that time. This was the same in other times.

This study presented the statistical analysis procedure and the appropriate cycle to improve TDLAT performance when measuring time series-based 2D temperature in idle mode of engine. To the best of our knowledge, this is the first attempt to present the criteria for cycle selection through the similarity analysis when based on signal stabilization technique. This signal stabilization technique was more effective in error reduction than the conventional polynomial reduction method when converting signals into absorption spectra.

However, limitations of this study include (*i*) it was only tested in idle mode of a particular engine, so additional experiments are required in operation mode with relatively large temperature variations; and (*ii*) in idle mode, the temperature variation in the peak area was uniformly measured only for that area, but it is also necessary to check the need for measurements in other areas.

To overcome these limitations, the following studies will be carried out in the future: (*i*) conduct a time series-based 2D temperature measurement study by applying the latest statistical techniques (i.e., linear Bayesian absorbance inference, Tikhonov regularization, and so on) to provide a fast and accurate case of reconstruction temperature calculation and (*ii*) typically, the reconstruction temperature is verified by TC measurements. The reliability of verification will depend on the TC measurement method, therefore studies on the TC verification methodology will also be conducted; and (*iii*) the size of the laser grid will affect the bias of reconstruction temperature. We would like to conduct a study on the selection of optimal grid sizes to minimize bias.



Figure 11. Comparison of T_{Thc} and T_{Rec} for 201 cycles.



Figure 12. 2D reconstruction temperature distributions at 201 cycles: (**a**) 1.0 s; (**b**) 2.0 s; (**c**) 3.0 s; (**d**) 4.0 s; (**e**) 5.0 s; (**f**) 6.0 s; (**g**) 7.0 s; (**h**) 8.0 s; and (**i**) 9.0 s.

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Appendix A

The Two-sample *t* test assumes a normality and equivalence. Since normality was satisfied, only the results of the equivalence test were summarized in Table A1. The equivalence is satisfied when p-value is more than 0.05.

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	Time (s)											
Cycles ⁻	1	2	3	4	5	6	7	8	9			
-	<i>P</i> -Value											
31	< 0.05	0.055	< 0.05	< 0.05	0.094	< 0.05	< 0.05	< 0.05	< 0.05			
51	< 0.05	< 0.05	0.175	< 0.05	0.062	0.105	< 0.05	< 0.05	0.205			
71	< 0.05	< 0.05	0.127	< 0.05	0.117	< 0.05	< 0.05	< 0.05	0.071			
91	0.071	< 0.05	0.109	< 0.05	0.265	0.073	0.071	< 0.05	0.099			
111	0.160	0.053	0.142	< 0.05	0.426	0.162	0.392	0.060	0.207			
151	0.099	0.213	0.218	< 0.05	0.401	0.706	0.106	< 0.05	0.143			
201	0.581	0.215	0.152	0.168	0.997	0.617	0.104	< 0.05	0.832			
251	0.899	0.148	0.115	0.280	0.645	0.688	0.388	0.302	0.678			
301	0.349	0.276	0.066	0.355	0.466	0.173	0.801	0.635	0.537			
351	0.398	0.285	0.220	0.181	0.614	0.271	0.660	0.758	0.391			
401	< 0.05	0.248	0.153	0.358	0.499	< 0.05	0.529	0.565	0.062			
451	0.054	0.243	0.284	0.227	0.348	0.064	0.358	0.867	0.125			
501	< 0.05	0.427	0.379	0.562	0.332	< 0.05	0.114	0.127	< 0.05			
601	< 0.05	0.548	0.794	0.854	0.387	< 0.05	< 0.05	< 0.05	< 0.05			
701	< 0.05	0.247	0.832	0.833	0.281	< 0.05	< 0.05	< 0.05	< 0.05			
801	< 0.05	0.413	0.607	0.565	0.231	< 0.05	< 0.05	< 0.05	< 0.05			
901	< 0.05	0.352	0.729	0.441	0.210	< 0.05	< 0.05	< 0.05	< 0.05			
1001	< 0.05	0.687	0.343	0.383	0.103	< 0.05	< 0.05	< 0.05	< 0.05			

Table A1. Result of the equal variance test between T_{Thc} and T_{Rec} for 18 candidate cycles.

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