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Elastic and inelastic light scattering from single bacterial spores in an optical trap allows the monitoring of spore germination dynamics

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Abstract

Raman scattering spectroscopy and elastic light scattering intensity (ESLI) were used to simultaneously measure levels of Ca-dipicolinic acid (CaDPA) and changes in spore morphology and refractive index during germination of individual *B. subtilis* spores with and without the two redundant enzymes (CLEs), CwlJ and SleB, that degrade spores' peptidoglycan cortex. Conclusions from these measurements include: 1) CaDPA release from individual wild-type germinating spores was biphasic; in a first heterogeneous slow phase, T_{lag} , CaDPA levels decreased $\sim 15\%$ and in the second phase ending at $T_{release}$, remaining CaDPA was released rapidly; 2) in L-alanine germination of wild-type spores and spores lacking SleB: a) the ESLI rose ~ 2 -fold shortly before T_{lag} at T_1 ; b) following T_{lag} , the ESLI again rose ~ 2 -fold at T_2 when CaDPA levels had decreased $\sim 50\%$; and c) the ESLI reached its maximum value at $\sim T_{release}$ and then decreased; 3) in CaDPA germination of wild-type spores: a) T_{lag} increased and the first increase in ESLI occurred well before T_{lag} , consistent with different pathways for CaDPA and L-alanine germination; b) at $T_{release}$ the ESLI again reached its maximum value; 4) in L-alanine germination of spores lacking both CLEs and unable to degrade their cortex, the time $\Delta T_{release}$ ($T_{release} - T_{lag}$) for excretion of $\geq 75\%$ of CaDPA was ~ 15 -fold higher than that for wild-type or *sleB* spores; and 5) spores lacking only CwlJ exhibited a similar, but not identical ESLI pattern during L-alanine germination to that seen with *cwlJ sleB* spores, and the high value for $\Delta T_{release}$.

INTRODUCTION

Optical trapping, also called optical tweezers, can capture and manipulate biological particles using low-power near-infrared laser beams and has allowed extensive study of single cells and molecules (1–5). Here we report use of simultaneous elastic and inelastic light scattering to monitor the germination of single bacterial spores in an optical trap following addition of various germinants

When a trapped cell is illuminated with the same or a second laser beam, it scatters light in all directions. The scattered light contains both elastic and inelastic components; the elastic scattered light has the same frequency as the illuminating laser beam and the inelastic scattered light has shifted frequencies. Elastic scattering light intensity (ESLI) can provide

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morphological information about a cell's size, shape and refractive index (6–7). Analysis of the inelastic scattered light, or Raman spectroscopy, can provide information about cells' molecular composition, since Raman scattering intensity is dependent on the amount of any particular molecular component (8–10). Consequently, simultaneous measurement of elastic and inelastic light scattering may provide valuable information on individual cells.

Spores of bacteria of *Bacillus* species are metabolically dormant, very resistant to a variety of harsh conditions, and can survive for many years (11–12). However, spores can rapidly return to active growth through germination followed by outgrowth (13). Germination is normally triggered by specific nutrients, although can also be initiated by some non-nutrient agents. Nutrient germinants bind to specific receptors located in the spore's inner membrane, triggering the release of spore small molecules, notably the large depot (~10% spore dry wt) of pyridine-2,6-dicarboxylic acid (dipicolinic acid (DPA)) (13–14). DPA is located exclusively in the spore's central region or core as a 1:1 chelate with divalent cations, predominantly Ca^{2+} (CaDPA). Release of small molecules and their replacement by water comprise Stage I of germination, and Stage I events, in particular CaDPA release, trigger hydrolysis of the spore's peptidoglycan (PG) cortex by either of two redundant cortex-lytic enzymes (CLEs), CwlJ and SleB, in *B. subtilis* spores. Cortex hydrolysis then allows swelling of the spore core and further water uptake, resulting in a core water content similar to that in growing cells. This full core rehydration completes Stage II of germination, and allows metabolism and macromolecular synthesis to begin (13).

The kinetics of germination of a spore population is usually monitored by measuring the optical density at 600 nm (OD_{600}) of spore cultures, which falls ~60% upon completion of Stage II of germination (13), or by measuring DPA release (15–18). However, the kinetics of germination of individual spores cannot be determined from these population measurements due to significant heterogeneity in the germination of individual spores (19–21). Kinetic germination would greatly change the amount CaDPA present in or release from the spores, which was recently used as a biomarker for the rapid detection of *Bacillus* spores by FAST coherent anti-Stokes Raman spectroscopy (22,23).

The kinetics of germination of single bacterial spores has been monitored by phase contrast microscopy (21), as a phase-bright dormant spore becomes phase-dark upon germination due to the decrease in the core's refractive index accompanying germination. However, the full decrease in the core's refractive index is due to CaDPA release as well as to full core rehydration (13). The kinetics of CaDPA release during germination of single spores has also been followed by Raman (19) and surface enhanced Raman spectroscopy (24). In particular, laser tweezers Raman spectroscopy (LTRS) has demonstrated significant heterogeneity in the kinetics of CaDPA release during germination of individual spores (19). However, there have been no simultaneous measurements of both morphological changes in and CaDPA release from single germinating spores, which could give precise ordering of various events during germination.

In this work, we report measurements of the ESLI and Raman scattering from single *B. subtilis* spores with and without CLEs following addition of various germinants using the experimental setup shown in Fig. 1. The spore was held in an optical trap and analysis of its Raman scattering allowed determination of the kinetics of CaDPA release, while ESLI gave information about changes in spore morphology and refractive index. These analyses have provided new insight into the process of bacterial spore germination.

MATERIALS AND METHODS

Optical trap, Raman spectroscopy and ESLI

The optical trap and Raman spectroscopy setup was as described (10,19). As shown in Fig. 1, a germinating spore in liquid is trapped in the focus of the laser beam about 10 μm above the coverslip. The backwards inelastic Raman scattering light from the trapped spore passes through the dichroic mirror and a notch filter, and is focused onto the entrance slit of a spectrograph and detected by a charge-coupled detector (Symphony CCD, Jobin Yvon). Raman spectra were recorded from 500 to 1800 cm^{-1} with a spectral resolution of $\sim 6 \text{ cm}^{-1}$. The backwards elastic scattering light from the trapped spore is reflected by a beam splitter (BS), passes through a band-pass filter, and is detected by an avalanche photodiode (APD) (Hamamatsu S2383) and recorded with a personal computer (PC) equipped with a data acquisition card (National Instruments, Inc. PCI-6024E). The sensitive area of the APD was much larger than the image of trapped spore so that the total ESLI was detected. A video camera was used to record the time-lapse images of the trapped spore (not shown in Fig. 1).

Strains, spore preparation and storage

B. subtilis strains used in this work were isogenic and were: PS832, a prototrophic laboratory derivative of strain 168; PS533 (25) carrying plasmid pUB110 that encodes resistance to kanamycin (10 $\mu\text{g}/\text{ml}$); strain FB113 (26) lacking CwlJ and SleB; and strains FB111 and FB112 (26) lacking CwlJ or SleB, respectively. Spores of *B. subtilis* strains were prepared at 37°C on 2xSG medium agar plates and purified and stored as described (27). All spore preparations used in this work were free (>98%) of growing or sporulating cells and germinated spores as determined by phase contrast microscopy.

Spore germination

Prior to germination with L-alanine, *B. subtilis* spores in water were heat activated for 30 min at 70°C and then cooled on ice; heat activation was not needed for CaDPA germination. *B. subtilis* spores were germinated in a microscope sample holder kept either at 37°C with 10 mM L-alanine in 25 mM Tris-HCl buffer (pH 7.4), or at 25°C with 60 mM CaDPA (1:1 mixture of 120 mM CaCl_2 and 120 mM DPA, made pH 7.5 with Tris base).

Monitoring germination dynamics of individual spores

After adding spores to pre-heated buffer plus germinants, a single spore was trapped at the focus of the near-infrared laser beam with a power of 3 mW. Raman spectra of the trapped spore were acquired with a CCD acquisition time of 45 s until spore germination as monitored by DPA release was complete or for 60 min. The intensities and bright-field images of elastic scattering light from the spore were recorded by the APD with a resolution time of 15 ms and a video camera, respectively. This procedure was repeated for another individual spore following loading of a new spore sample. The CaDPA level in individual spores was determined from intensities of the Raman spectral band at 1017 cm^{-1} (10,19), normalized to the CaDPA level at time zero, and plotted as a function of the germination time. The effect of the low-power optical trap ($\sim 3 \text{ mW}$) on spore germination was minimal due to the low absorption in the near-IR region (19).

Spore germination parameters determined from these data (Fig. 2) were: a) T_{lag} - the time between addition of a germinant and initiation of rapid Ca-DPA release; as shown in Fig. 2B, the rate of the CaDPA release during L-alanine germination was initially slow, but became rapid as remaining CaDPA was released; the intersection of the slow release slope with the rapid release slope gives T_{lag} (19). The percentage of initial CaDPA remaining in the spore at T_{lag} was determined by the measured amount of CaDPA at the time point nearest to T_{lag} , with

a precision of the acquisition time of 45 s; b) T_1 – the time of initiation of the first rapid rise in ESLI; c) T_2 – the time of initiation of the second rapid rise in ESLI; d) T_{release} – the time when release of CaDPA is complete, determined by the intersection of the rapid release slope with the zero level of CaDPA; and e) $\Delta T_{\text{release}} = (T_{\text{release}} - T_{\text{lag}})$.

RESULTS

Germination of *B. subtilis* spores in L-alanine

Fig. 2A shows Raman spectra and images of elastic scattering light from a single wild-type *B. subtilis* spore germinating with L-alanine. These data indicate that: 1) peak heights of CaDPA specific bands at 824, 1017, 1395, and 1572 cm^{-1} fell slightly during T_{lag} (~17 min) following addition of L-alanine (and see below), and then decreased rapidly and essentially to zero by T_{release} (~20 min) as seen previously (19); however, intensities of Raman spectral bands from other spore components such as phenylalanine (1004 cm^{-1}) and the protein amide bond (1655 cm^{-1}) remained unchanged after CaDPA release; 2) images of elastic scattering light from the spore prior to 15 min (~ 2 min before T_{lag}) did not change; as seen in the 12 min image, the brightness of the elastic scattering pattern was relatively weak with only the spore core efficiently generating scattering light (note that the spore's total size can be determined from the dark edge in the 12 min image); 3) the brightness and size of the spore's strong elastic scattering pattern increased abruptly just prior to T_{lag} (Fig. 2A, shown at 16:10; and see below), and increased again at about T_{release} (Fig. 2A, shown at 20:50); 4) after T_{release} , the brightness of the elastic scattering pattern gradually decreased, although the overall size of the elastic scattering pattern was nearly unchanged. This behavior was seen in analyses of multiple individual spores, although values of T_{lag} and T_{release} varied considerably (see below).

Fig. 2B shows the intensity of the CaDPA-specific 1017- cm^{-1} band of this same single germinating spore normalized to its value at the first time of measurement; Following L-alanine addition, CaDPA release began slowly but became more rapid at T_{lag} . While most CaDPA was released between T_{lag} and T_{release} , ~20% was released prior to T_{lag} . Fig. 2C shows the total ESLI from this same spore following L-alanine addition. As indicated above, the ESLI remained low until just prior to T_{lag} , but ~2 min earlier (~15 min) the ESLI began to increase rapidly, rising ~2-fold by T_{lag} and remaining nearly constant prior to T_2 (~ 19.3 min), when the ESLI increased again to a maximum at T_{release} (~20.7 min) and then gradually decreased. The increases in the ESLI prior to, during and after the release of spore's CaDPA indicated that significant changes in either the state or the morphology of the spore are taking place at these times.

The other information given by the ESLI from a single spore during L-alanine germination was in the intensity fluctuation of the scattered light as reflected in the amplitude of the intensity curve (Fig. 2C). While the intensity fluctuation was low before T_1 , the fluctuation increased shortly after T_1 , and increased further after T_{release} . The increased fluctuation in ESLI may be due to increases in the Brownian motion of scattering centers in or on the spore, since the detected ESLI is the coherent superposition of the scattering light fields from the spore's various scattering centers.

Analysis of multiple individual wild-type spores germinating with L-alanine (Fig. 3a,b) revealed the same general features described above. Thus 10–20% of total CaDPA was released in T_{lag} , and the biggest variation in the kinetic parameters for CaDPA release between different spores was in values for T_{lag} (Table 1), as seen with germinating *B. thuringiensis* spores (19). However, while T_{lag} values varied considerably, the timing of T_{lag} , T_{release} , T_1 and T_2 relative to one another were identical, and the time for release of $\geq 75\%$ of spore CaDPA ($\Delta T_{\text{release}}$) was always 2–3 min (Fig. 3a; Table 1).

Germination of *B. subtilis* spores lacking CLEs

The changes in the ESLI during wild-type spore germination with L-alanine described above could be due to morphological changes accompanying cortex PG hydrolysis, to changes in the hydration of the spore core accompanying CaDPA release, or to both processes. To determine which of the latter are most important, we germinated *cwlJ sleB* spores that cannot degrade cortex PG during spore germination with L-alanine (26). The results (Fig. 4a,b) were striking as: i) the $\Delta T_{\text{release}}$ for CaDPA increased to 30–60 min; ii) most of the changes seen in ESLI with germinating wild-type spores were not seen, as the most notable change was a moderately slow (5–10 min) increase of ~0.5–2-fold in scattering intensity that began when ~25% of CaDPA had been released, and ended when ~75% had been released; after this point there were no further notable changes; and iii) the amplitude of the fluctuation in ESLI did, however, increase notably after the increase in intensity noted above.

To determine which CLE, CwlJ or SleB was responsible for increasing $\Delta T_{\text{release}}$ during L-alanine germination, we examined CaDPA release and ESLI from individual spores lacking only CwlJ or SleB (Table 2). While values of T_{lag} varied considerably between different spore preparations for reasons that remain unclear, $\Delta T_{\text{release}}$ values were 2–4 min for spores that contained both CwlJ and SleB or only CwlJ. In contrast, spores that lacked CwlJ, either alone or along with SleB, had $\Delta T_{\text{release}}$ values of ~42 min. Thus only CwlJ plays a role in accelerating rates of CaDPA release from *B. subtilis* spores germinating in L-alanine.

The parameters for changes in ESLI during L-alanine germination of spores lacking only SleB were also similar to those obtained with wild-type spores, when the longer T_{lag} value of the *sleB* spores was ignored (Table 2). However, during L-alanine germination of spores lacking only CwlJ there was only a single rapid increase in ESLI, as seen with germinating *cwlJ sleB* spores, although the increase in ESLI with *cwlJ* spores took place well before T_{lag} , while it was well after T_{lag} with germinating *cwlJ sleB* spores (Table 2).

Germination of *B. subtilis* spores in exogenous CaDPA

In addition to nutrient germinants, spores can also germinate with a number of non-nutrient agents, one of which is CaDPA itself, and the pathway for spore germination with CaDPA is different than that with nutrient germinants (13). Raman spectra from individual wild-type *B. subtilis* spores germinating with CaDPA showed that the peak heights of CaDPA specific bands at 824, 1017, 1395, and 1572 cm^{-1} slowly decreased prior to T_{lag} and then rapidly decreased to zero by T_{release} after subtraction of the signal due to external CaDPA (Fig. 5a; and data not shown). As with L-alanine germination, absolute values for T_{lag} and T_{release} in CaDPA germination were heterogeneous and intensities of Raman peaks from other spore components did not change even after full CaDPA release (Fig. 5a; Table 2; and data not shown).

In contrast to L-alanine germination in which T_1 , the time the ESLI first began to increase, was only slightly prior to T_{lag} , with CaDPA germination T_1 was well before T_{lag} (Fig. 5b). Following T_1 the ESLI increased slowly until leveling off at T_2 , again well before T_{release} (Table 2). Between T_{lag} and T_2 , the majority of spore's CaDPA was still in the spore's core (Fig. 5a), but the ESLI increased sharply ~2-fold beginning at T_2 , reaching a maximum level at T_{release} (Fig. 5b). After T_{release} , the ESLI gradually decreased, and as seen with L-alanine germination, the intensity fluctuation increased when the ESLI increased following T_2 , and the fluctuation increased further after T_{release} .

DISCUSSION

Two parameters have been measured during the germination of individual *B. subtilis* spores with the nutrient germinant L-alanine, CaDPA levels and ESLI. These analyses confirm

features of CaDPA release seen during wild-type *B. thuringiensis* spore germination with L-alanine (19) including: i) a significant T_{lag} between addition of a germinant and rapid CaDPA release; ii) a marked heterogeneity in T_{lag} between individual spores; and iii) release of the great majority of CaDPA in only a few min ($\Delta T_{release}$) as the spore's CaDPA level falls to zero at $T_{release}$. These features of CaDPA release were also seen in germination by exogenous CaDPA, which triggers germination by activating CwlJ (26). Notably, T_{lag} values were heterogeneous for CaDPA germination while $\Delta T_{release}$ times were comparable to those for L-alanine germination.

One novel finding in this work is that during nutrient and CaDPA germination of *B. subtilis* spores there was slow release of 15–20% of the spores' CaDPA in T_{lag} that preceded rapid release of the majority of CaDPA during $\Delta T_{release}$. Reexamination of data on CaDPA release from *B. thuringiensis* spores during nutrient germination (19) has also indicated that 10–20% of total CaDPA is released in T_{lag} during the germination of these spores (data not shown). The significance of this slow CaDPA release in T_{lag} is not known, but it is tempting to speculate that this is important in germination, perhaps by slowly elevating the core water content. A second novel finding was that values for $\Delta T_{release}$ were accelerated greatly by the CLE CwlJ, while SleB had no effect. A decrease in the rate of CaDPA release during nutrient germination of *B. subtilis* *cwlJ* spore populations has been reported (28). However, current work indicates that the effect of CwlJ is on $\Delta T_{release}$ not on T_{lag} and this effect of CwlJ has also been seen during nutrient germination of individual *B. megaterium* spores (29). Thus CwlJ action somehow accelerates CaDPA efflux across the spore's inner membrane during germination.

The ESLI analysis revealed some unexpected features of spore germination, as there were two large, rapid increases in ESLI during L-alanine germination of wild-type spores, one at T_1 just prior to T_{lag} and a second at T_2 during $\Delta T_{release}$; these two increases were then followed by a slow decline. The two changes in ESLI were also seen during CaDPA germination, but there was only one increase during *cwlJ* spore germination. Changes in ESLI could be due to changes in spore size, shape or refractive index (6,7). Overall spore shape, including that of the spore core, does not change noticeably during the complete process of spore germination (30), so changes in this parameter seem unlikely to change the ESLI observed in the current work. Similarly, although the volume encompassed by the spore core increases 2- to 3-fold upon completion of cortex lysis and core swelling, the size of the spore does not change, and the ESLI was obtained from the entire *B. subtilis* spore. This leaves changes in refractive index as the likely cause of changes in ESLI during spore germination. The physical origin of increases in ESLI could be an increase in the inhomogeneity of refractive index inside the spore or on the spore's surface. This effect generates more scattering centers so that more incident photons are scattered from cellular compartments that have a higher refractive index than their surroundings, even though the overall refractive index of the spore is not increased. Prior to the addition of nutrient or non-nutrient germinants, the refractive index is relatively uniformly distributed in the spore, except for the high value in the core due to its extremely high level of CaDPA. Thus, the observed ESLI is relatively weak from a dormant spore (Fig. 2A). However, dramatic changes take place in the refractive index of the spore core during germination, first as CaDPA is released and replaced by water, and then as cortex lysis and core swelling proceed to completion. The key question is what causes the changes in ESLI seen during spore germination in L-alanine. There appear to be a number of major possibilities: 1) abrupt mixing of water in the spore core resulting in solubilization of some of the core's CaDPA; 2) full release of CaDPA resulting in a drastic fall in core refractive index; and 3) full core hydration due to cortex lysis, core swelling and water uptake. Surprisingly, the first rapid increase in ESLI generally took place before T_{lag} , when only small amounts of CaDPA had been released, while the second increase took place midway in $\Delta T_{release}$, when ~ 50% of CaDPA had been released. These same increases in ESLI were also seen in CaDPA germination, but while the increase at T_2 was at the same time relative to T_{lag} as in L-alanine germination, T_1 was well

before T_{lag} , and well before release of much CaDPA. In contrast to the two increases in ESLI seen during germination of wild-type spores or those lacking only SleB, spores lacking CwlJ or CwlJ and SleB exhibited only one increase. This increase was after T_{lag} with *cwlJ sleB* spores, but before T_{lag} with *cwlJ* spores.

As noted above, changes in ESLI due to alterations in refractive index could be due to abrupt changes in the heterogeneity of the refractive index in some spore compartment – most likely the core. Thus the first change could be due to the initiation of mixing of water in the spore core beginning the process of solubilizing the core's CaDPA that likely exists primarily in an insoluble form. The second abrupt change in ESLI could then be due to a second rapid decrease in core refractive index accompanying the swelling of the spore core due to cortex hydrolysis. CwlJ and SleB are located primarily on the outer and inner edges of the cortex, respectively (13), and thus we presume that CwlJ hydrolyses cortical PG from the outside in, while SleB works from the inside out. This might result in an abrupt jump in core volume when the two waves of cortex hydrolysis meet. However, when SleB is the only CLE present, the core may swell slowly and smoothly. The slow decrease in ESLI seen after its final rise could then be due to a slow decrease in the heterogeneity of the core's refractive index due to complete mixing of core components following full core expansion. Unfortunately, we do not yet know precisely when cortex hydrolysis actually takes place relative to CaDPA release during nutrient germination, whether CwlJ and SleB do indeed work only from the outside in and the inside out, respectively, and how rapidly the core expands when the cortex is hydrolyzed, since this expansion requires remodeling of the germ cell wall to accommodate the core's increased volume. We also do not know the rate of diffusion of various core components in various stages of spore germination, although a small, soluble protein is essentially immobile in dormant and Stage I germinated spores after CaDPA has been released, and becomes mobile upon completion of germination (31). We also do not know the meaning of the increase in fluctuation of the ESLI as germination proceeds. However, most of this increase took place only after the first rise in ESLI. Further complicating analysis of events in spore germination is that recent work has indicated that muropeptides (MP) produced by PG degradation, perhaps even by hydrolysis of the spore cortex, can also trigger germination, and by a pathway distinct from the nutrient germinant receptor pathway (32). Thus triggering cortex hydrolysis via the germinant receptor pathway in an individual spore could result in further stimulation of germination of that spore via the MP germination pathway. However, the details of the MP germination pathway have not been determined.

In conclusion, Raman scattering spectroscopy and ESLI have been used to simultaneously measure levels of CaDPA and changes in spore morphology and refractive index during germination of individual *Bacillus* spores. These analyses have provided new insight into the process of bacterial spore germination.

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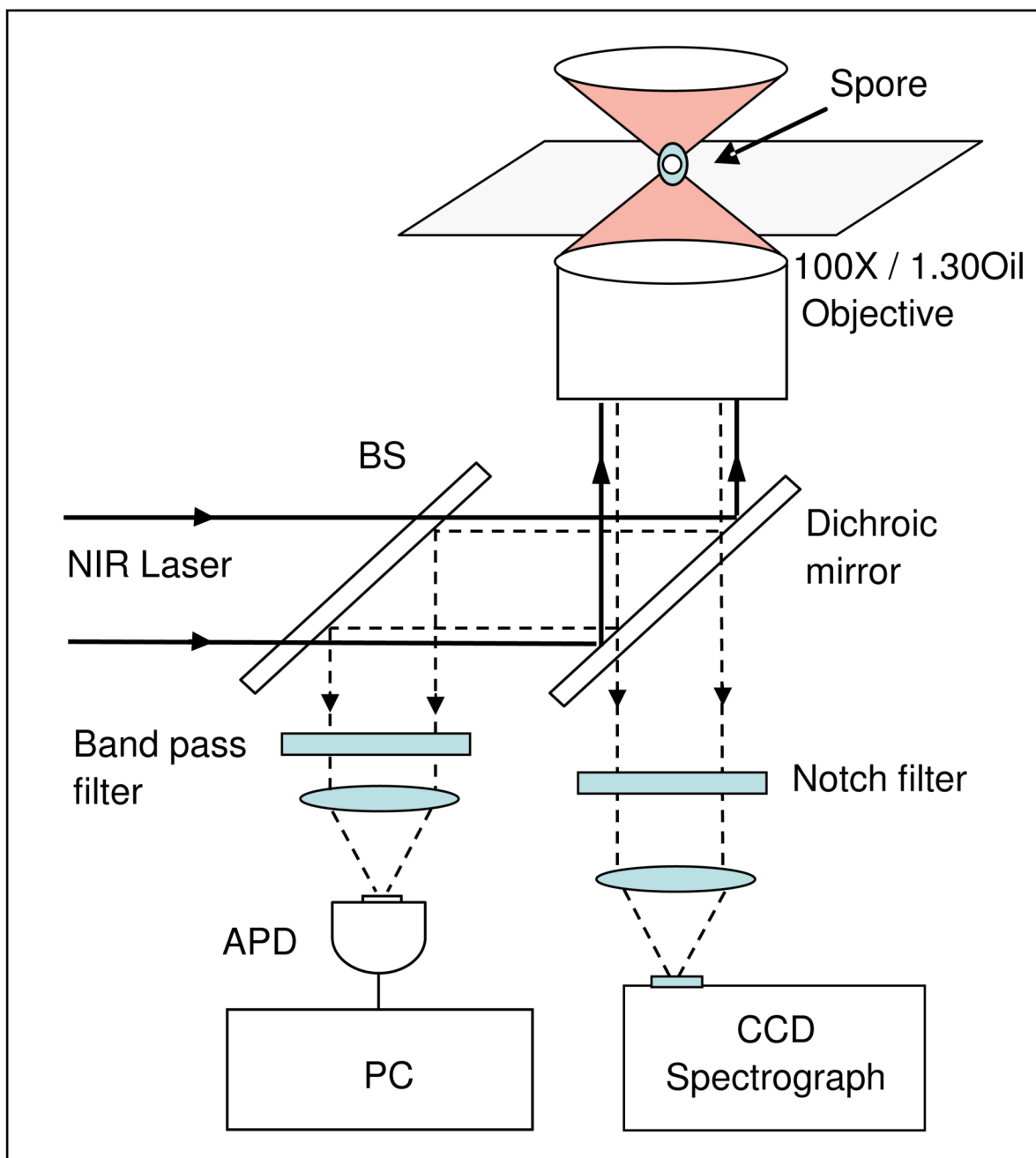


Figure 1. Experimental setup for analysis of elastic and inelastic light scattering from a single optically trapped spore. A near-infrared (NIR) laser beam at 785 nm is introduced in an inverted microscope (Nikon TE2000 DIC) equipped with an objective by a dichroic mirror to form a single-beam optical trap. A germinating spore in liquid is trapped in the focus of the laser beam about 10 μm above the coverslip.

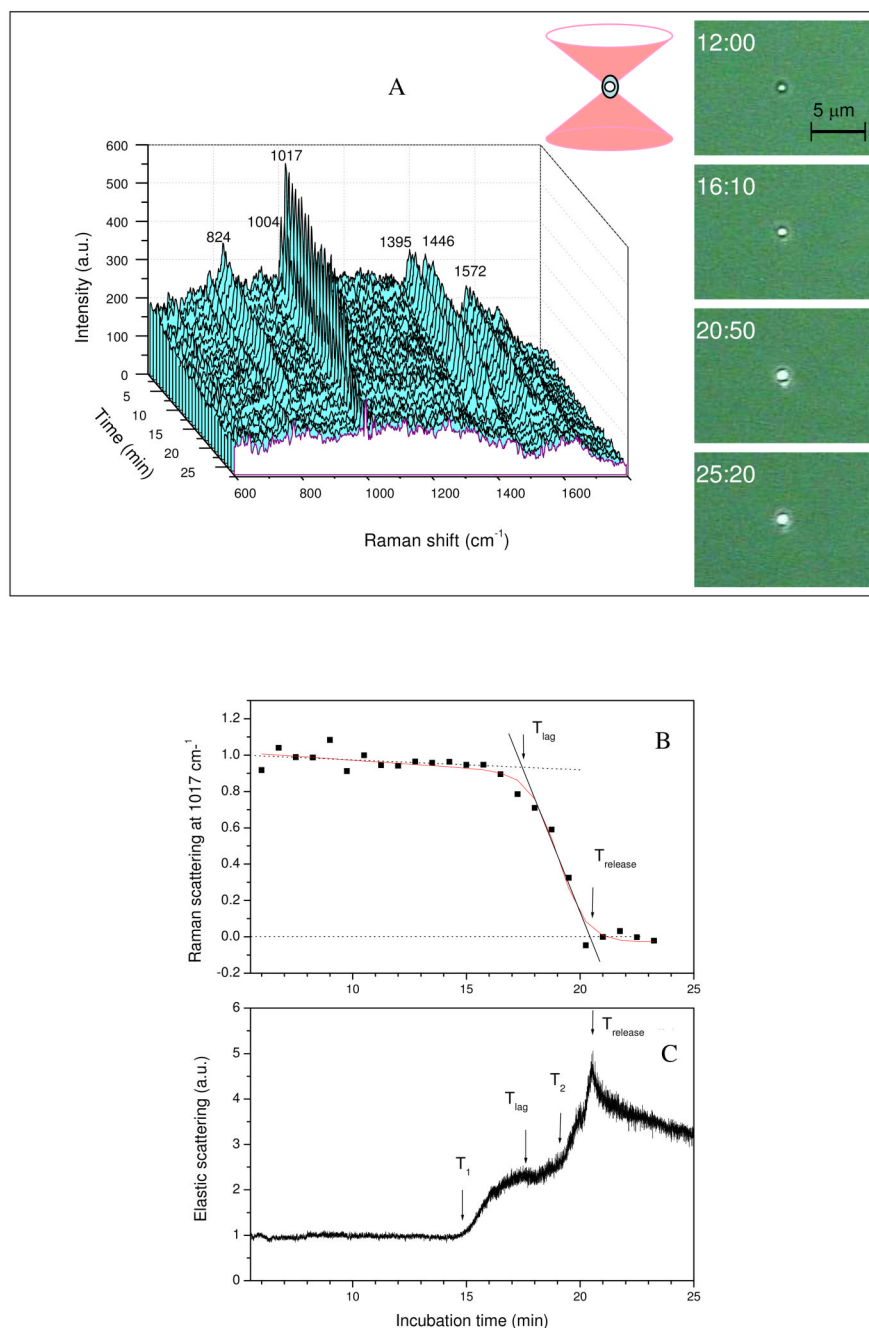


Figure 2.

Figure 2A–C. A) Raman spectra and images, B) intensities of the CaDPA-specific Raman spectral band, and C) ESLI from a *B. subtilis* wild-type spore germinating in L-alanine in an optical trap. The Raman spectra, images and ESLI and the intensity of the CaDPA-specific 1017- cm^{-1} Raman band from a *B. subtilis* PS533 (wild-type) spore germinating in L-alanine were determined as described in Methods. The red solid line in Fig. 2B was a mathematical fit with a modified sigmoidal curve. The Raman band intensity and ESLI at each time point were normalized to values at the first time of measurement. For all measurements, $\lambda_{\text{exc}} = 785 \text{ nm}$, $P_{\text{exc}} = 3 \text{ mW}$, and the acquisition time was 45 s for Raman scattering light and 15 ms for ESLI. The dark ring in the image at 12 min in A) gives the size of the trapped spore.

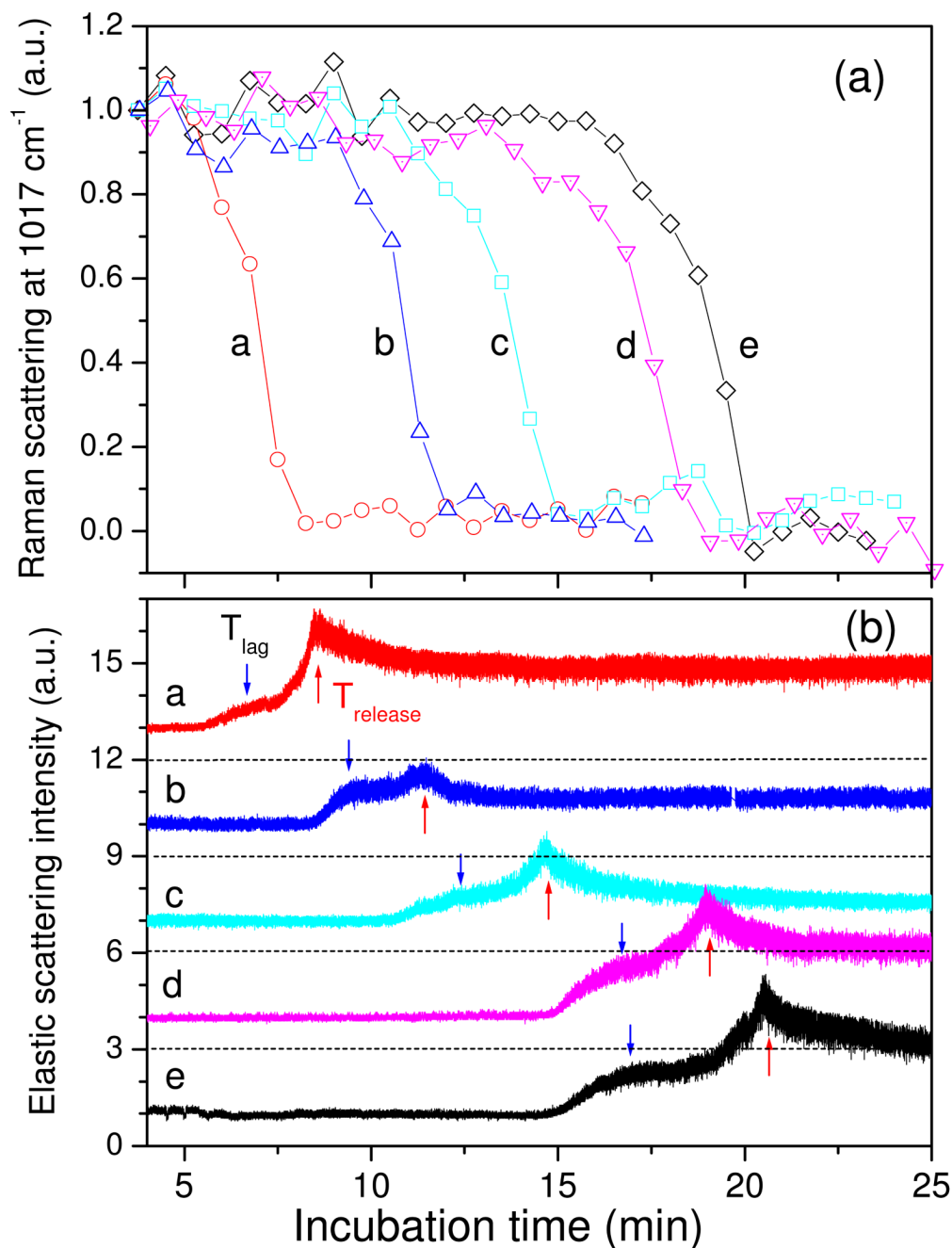


Figure 3.

Figure 3a,b. CaDPA release and ESLI from five wild-type *B. subtilis* spores during L-alanine germination. a) The relative intensities of the 1017-cm⁻¹ CaDPA Raman band from five individual *B. subtilis* PS533 (wild-type) spores (a-e) germinating with L-alanine were determined as described in Methods and values are given relative to those at the beginning of the experiment. b) The relative ESLI from the five spores described in a) were determined as described in Methods, and were normalized to the first values measured. Normalized ESLI for spores a-d were shifted upwards by 3 on the y-axis for display purposes. The dashed lines at 3, 6, 9, and 12 scattering intensities are the levels of zero ESLI for spores d, c, b and a,

respectively, and the blue and red arrows indicate the T_{lag} and $T_{release}$ times, respectively, for each spore.

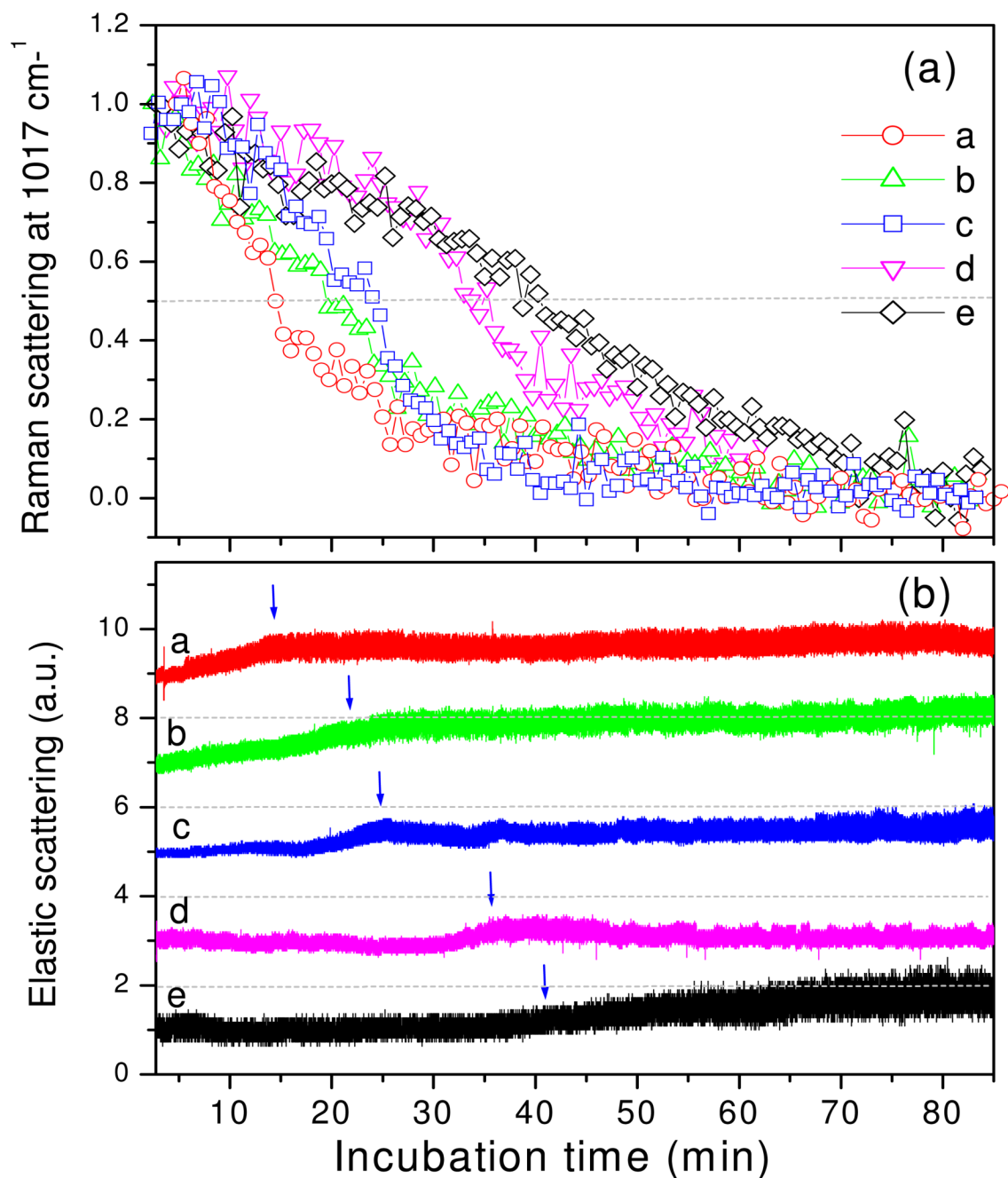


Figure 4.

Figure 4a,b. CaDPA release and ESLI from *B. subtilis* spores that cannot degrade their cortex during L-alanine germination. a) The relative intensities of the 1017-cm⁻¹ CaDPA band from five *B. subtilis* FB113 (*cwlJ sleB*) spores that do not degrade their cortex during germination with L-alanine were determined as described in Methods. b) The ESLI from the five spores germinating in a) were determined as described in Methods. These latter values were normalized to values obtained in the first measurement, and normalized values were shifted by 2 on the y-axis for display purposes. The dashed lines at 2, 4, 6 and 8 are the levels of zero ESLI for spores d, c, b and a, respectively, and the vertical blue arrows indicate the time at which the DPA level of individual spores was reduced by 50%.

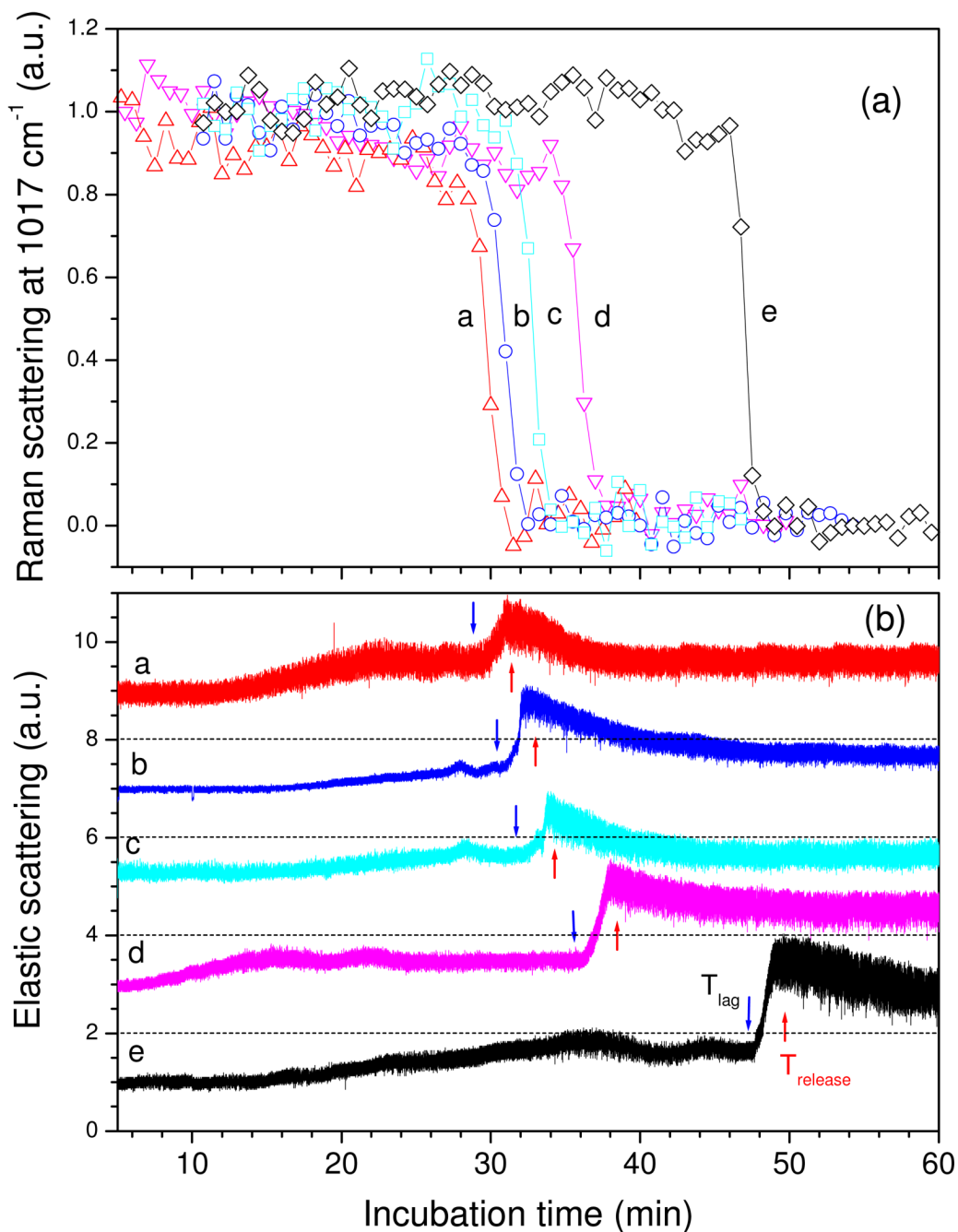


Figure 5.

Figure 5a,b. CaDPA release and ESLI from five *B. subtilis* spores germinating with CaDPA. a) The intensities of the CaDPA-specific 1017-cm⁻¹ Raman band from five *B. subtilis* PS533 (wild-type) spores (a-e) germinating with CaDPA were determined as described in Methods; the signal intensity due to the CaDPA added to trigger germination has been subtracted. b) The ESLI normalized to the values of the first measurement from the five spores germinating with CaDPA in a) were determined as described in Methods. The normalized ESLI values were shifted by 2 on the y-axis for display purposes. The dashed dark lines at 2, 4, 6, and 8 are the zero levels of ESLI for spores d, c, b and a, respectively, and the blue and red vertical arrows indicate the T_{lag} and T_{release} times, respectively, for the individual spores.

Table 1

Values of T_1 , T_{lag} , T_2 , and $T_{release}$ and CaDPA remaining in spores during germination of 5 wild-type spores with L-alanine. Values are from Fig. 3a,b, and values in parentheses are the percentage of initial CaDPA remaining in the spore at these times.

Spore #	T_1 - min	T_{lag} - min	T_2 - min	$T_{release}$ - min
a	5.4 (90)	6.4 (71)	7.6 (17)	8.5 (0.8)
b	8.3 (94)	9.3 (90)	10.7 (60)	11.6 (13)
c	10.5 (99)	11.3 (82)	13.5 (60)	14.8 (11)
d	14.7 (87)	16.5 (69)	17.5 (60)	19.1 (0.1)
e	14.8 (99)	17.0 (80)	19.3 (40)	20.7 (0.1)
Average	10.7 (94)	12.0 (78)	13.4 (48)	14.8 (5)

Table 2

Values of T_{lag} , $\Delta T_{release}$, T_1 , and T_2 for individual spores germinating in L-alanine or CaDPA. 10 spores of strains PS832, FB111, FB112, and FB113 were germinated with L-alanine and 5 spores of PS533 were germinated with L-alanine or CaDPA as described in Fig. 3–5, and values for T_{lag} , $\Delta T_{release}$, T_1 , and T_2 were determined as described in Methods. Values in parentheses are the percentage of initial Ca-DPA remaining in spores at T_{lag} .

Strain	T_{lag} - min	$\Delta T_{release}$ - min	T_1 - min	T_2 - min
PS533 (wt) in L-alanine	12.0 ± 4.5 (78.0)	2.7 ± 0.6	10.7 ± 4.1	13.4 ± 4.8
PS832 (wt) in L-alanine	6.3 ± 1.7 (82.3)	3.7 ± 0.8	4.6 ± 2.6	8.0 ± 2.3
PS533 (wt) in CaDPA	33.9 ± 7.0 (87.8)	2.3 ± 0.2	12.7 ± 4.4	35.1 ± 7.4
FB113 (<i>cwlJ sleB</i>) in L-alanine	9.5 ± 3.2 (89.5)	42.9 ± 13.3	21.0 ± 13.0	Not observed
FB111 (<i>cwlJ</i>) in L-alanine	33.5 ± 5.8 (78.9)	40.7 ± 8.6	24.1 ± 2.4	Not observed
FB112 (<i>sleB</i>) in L-alanine	36.7 ± 9.8 (68.1)	3.6 ± 2.1	34.4 ± 8.9	38.2 ± 9.9