Laser-stimulated fluorescence reveals unseen details in fossils from the Solnhofen Limestone (Upper Jurassic, Bavaria, Germany)

Luke Barlow (✉ up812045@myport.ac.uk)  
University of Portsmouth  https://orcid.org/0000-0002-0867-649X

Michael Pittman  
University of Hong Kong  https://orcid.org/0000-0002-6149-3078

David Martill  
University of Portsmouth

Thomas Kaye  
Foundation for Scientific Advancement  https://orcid.org/0000-0001-7996-618X

Anthony Butcher

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Abstract

Laser-Stimulated Fluorescence (LSF) has seen increased use in palaeontological investigations in recent years. The method uses the high flux of laser light to reveal details sometimes missed by ultraviolet (UV) and optical wavelengths. In this study, we compare the results of LSF with UV on a range of fossils from the Upper Jurassic Solnhofen Limestone Konservat-Lagerstätte of Bavaria, Germany. The methodology follows previous protocols with modifications made to enhance laser beam intensity. Our experiments show the value of LSF in revealing shallow subsurface detail of specimens, previously not widely applied to Solnhofen fossils. In particular, fossil decapods from the Solnhofen Limestone reveal full body outlines, even under the matrix, along with details of segmentation within the appendages such as limbs and antennae. The results indicate that LSF can be used on both vertebrate and invertebrate fossils and may surpass the information provided by traditional UV methods in some specimens.

Introduction

Since the introduction of ultraviolet (UV) fluorescence for the analysis of fossils from the Upper Jurassic Solnhofen Limestone lithographic limestones of Germany in the early 20th century (Miethe & Born 1928), the technique has been increasingly used in the analysis of exceptionally preserved fossils from this famous fossil Lagerstätte. Fossils from a wide range of phyla have been studied using UV including decapods (Schwiegert 2011), ammonites (Keupp 2007), fish (Tischlinger & Arratia 2013), pterosaurs (Frey et al. 2003) and dinosaurs (Göhlich & Chiappe 2006; Rauhut et al. 2012) among others. More recently, laser light techniques have been applied to fossils to induce fluorescence (Kaye et al. 2010), especially on ‘iconic’ fossil vertebrates from sites of exceptional preservation like Anchiornis, Confuciusornis, Jianianhualong, Psittacosaurus and Sapeornis (Vinther et al. 2016; Mayr et al. 2016; Xu et al. 2017; Kaye et al. 2019a,b; Pittman & Xu 2020; Serrano et al. 2020), including the first discovered fossil feather which is from the Solnhofen Limestone (Kaye et al. 2019a). Fossils from the lithographic limestone horizons of the Solnhofen Limestone have been widely studied using UV and thus provides an ideal opportunity to compare the two fluorescent techniques on a range of fossils including cephalopods, decapods, and small vertebrates (Hess 1999; Zhou & Wang 2010) (Fig. 1). The Solnhofen Limestone is famous for its well-bedded, ultrafine-grained lithographic limestones (often called plattenkalk) and referred to as such herein) that formed in the calm basins of the Solnhofen lagoons on the northern margin of the Tethys Ocean (Viohl 1998; Munnecke et al. 2008). The palaeoenvironment represented by these limestones is a closed lagoonal system with high evaporation rates leading to a stratified water column with anoxic bottom waters largely devoid of macroorganisms (Viohl 1998). Occasional mixing through storms brought the toxic water to the aerated surface zone leading to the mass mortality of nektonic organisms (Pan et al. 2019). These organisms often became exceptionally well preserved due to a lack of scavenging, bacterial sealing, and rapid burial (Wellnhofer 2009). Here we compare images of Solnhofen fossils under LSF and UV and evaluate the use of LSF on fossils in this important Konservat-Lagerstätte.
Methods

The specimens used in this study labelled LB 1–13, abbreviated from the initials of the primary author, were collected during a series of field visits to the Solnhofen region over twenty years and are accessioned in the collection of the School of the Environment, Geography and Geosciences (SEGG), University of Portsmouth. Additional specimens from the Staatliche naturwissenschaftliche Sammlungen Bayerns, Bayerische Staatssammlung für Paläontologie und Geologie (SNSB-BSPG) were studied by MP and TGK during a visit to the Museum für Naturkunde (Kaye et al. 2019a) (Figs 13–16). The specimens used are based on availability and represent some of the main groups found in the Solnhofen Limestones.

Photography

Photographs were taken in a blacked-out room to avoid natural light contamination. An LED lamp illuminated the specimen obliquely (c. 45 degrees) or directly for the white light photographs.

The method of Laser-Stimulated Fluorescence modified from Kaye et al. (2015) used an MGL-III–532–1~300mW green diode-pumped solid-state (DPSS) laser with a PSU-III-LCD power supply with a set output of 85mW. Alterations to the method included mounting the laser onto a camera track where only a trucking motion was permitted, so the 532 nm green laser moved through the x-axis to scan the specimen whilst maintaining the same perpendicular relationship (Fig. 2). Through substituting trucking for the panning motion in previous publications, the laser module maintains a constant distance from the specimen and therefore a constant beam intensity across the entire fossil. For Figs 13–16 the same methodology was used in Kaye et al. 2019a with an abbreviated method stated here. A 1W 405nm blue laser was used to induce fluorescence and a Nikon DSLR was used to take the photographs with a 425nm blocking filter. Post-processing (equalisation, saturation, and colour balance) was then performed in Photoshop CS6.

The method of ultraviolet fluorescence consisted of a 365 nm lamp as used by Tischlinger & Frey (2002) with the specimen illuminated as close as possible.

The long-exposures for each image under all three lighting regimes were taken using a Nikon D5300 DSLR camera mounted on a tripod with a 2-second self-timer setting that prevents camera movement from affecting the image. Aperture priority mode controlled the length of exposure following the method described by Eklund et al. (2018). They suggested a 10 second exposure for UV and the ISO was adjusted accordingly. Using a low ISO prevents grainy photos and with a small aperture, other light sources are prevented from contaminating the image. 30 second exposures were used for imaging under LSF with the ISO left at 100. The position and distance of the specimen from the camera remained constant for each method so that LSF, white light, and UV images could be collected efficiently. This lighting sequence allowed the O56 blocking filter that prevents camera over-saturation with LSF to be applied effectively. This filter was removed for white light and UV photography, as although the filter can pick up UV
fluorescence, the fluorescence is clearer without a blocking filter. Due to the flattened nature of the fossils within Solnhofen laminites, repetitive photography and photo stacking techniques were unnecessary.

**Health and Safety**

It is important that when using LSF or UV techniques, appropriate safety precautions are observed. The green laser and UV lamp require laser and UV blocking goggles during operation and these methods were conducted in a locked room with a suitable exterior notice to prevent people from being harmed. Humans are most sensitive to green laser light (Galang *et al.* 2010) and by following these guidelines, the operators and others can be safeguarded. When using UV, 10-minute breaks were taken every 10–30 minutes to prevent eye damage and headaches (Tischlinger & Arratia 2013). Since pale colours fluoresce under UV light, dark clothing was worn while conducting scans.

**Results**

**Cephalopoda**

Cephalopods in the form of ammonites, belemnites and teuthoids occur frequently in the Solnhofen Limestone and are sometimes exceptionally well preserved (Fuchs *et al.* 2015). Many have been reported with aspects of their soft tissues preserved, including impressions of tentacles with hooklets in belemnites and teuthoids (Klug *et al.* 2016), the musculature of the mantle in teuthoids (Klug *et al.* 2015) and the siphuncle and pellicula in ammonites (Keupp 2007). Although ammonites make up a large portion of the fossils from the Solnhofen plattenkalks, they are often poorly preserved due to the aragonitic composition of the shell which is readily dissolved during diagenesis (Seilacher *et al.* 1976). This dissolution leaves behind an external or composite mould in the matrix, occasionally with the original outline and the calcitic aptychi in the body chamber (Keupp 2007).

Although rarely preserved, the original shell can be observed once replaced by calcite in the phragmacone (Fig. 4) and the body chamber (Arratia *et al.* 2015). The body chamber is present within the halo of the dissolved shell and can be enhanced under UV and LSF (Fig. 6). The siphuncle, pellicula and other non-mineralised elements are often phosphatised in Solnhofen ammonites and these may fluoresce more intensely than the remaining shell (Keupp 2007). UV fluorescence displays colour differences on these ammonites (Fig. 6C), but the contrast between non-mineralised parts and the surrounding shell is lacking, when compared with the LSF image of specimen LB 4 (Fig. 6D). It appears that the raised darker areas on isolated aptychi (LB 1) (Fig. 3) are the thick spongy layer on the inside of the aptychus underlying the thinner crenulated outer layer, rather than soft tissues following Lehmann (1976).

*Lumbricaria* Goldfuss, 1831, a coprolite attributed to ammonites (Janicke 1970) lies on the same slab as an ammonite (Fig. 6) and appears to contain aptychi of a smaller ammonite. *Plesiotethis prisca* (Rüppell, 1829), a squid from the Solnhofen Limestone with a central rachis that is revealed in its entirety
and fluoresces at two distinct levels under LSF (Fig. 7D). This rachis runs along the centre of the visible body outline, although no soft tissues of the mantle are present (Fig. 7).

**Decapoda**

Decapod crustacea occur frequently in the Solnhofen plattenkalks and are often well preserved as fully articulated individuals, even in the larval stage (Haug *et al.* 2008; Winkler 2014). The fossils are valuable for understanding arthropod ontogeny, with specimens that display growth cycles only known from a few other Lagerstätten (J. T. Haug *et al.* 2009). Numerous studies have been carried out using UV fluorescence, revealing unseen details to allow distinctions between taxa or increasing the specimen-matrix contrast (Haug *et al.* 2011; Winkler 2012; Audo *et al.* 2014). The decapods lend themselves to techniques using fluorescence as they hold autofluorescent compounds within their exoskeleton (Haug *et al.* 2011; Glenn *et al.* 2013).

The preservation of the fossils studied (LB 6–9), vary from isolated appendages to complete articulated examples. A single walking leg attributed to *Aeger tipularius* von Schlotheim, 1822 (Fig. 8) is near complete with only a few setae missing from the exposed surface. UV fluorescence enhances the specimen-matrix contrast but fails to fluoresce below the matrix where the complete setae are revealed through LSF (Fig. 8).

Anatomical details of complete specimens of *Cycleryon propinquus* von Schlotheim, 1822 (Fig. 9) and *Antrimpos speciosus* Münster, 1839 (Figs 10, 11) are enhanced through fluorescent techniques as previous studies demonstrate (Schweigert & Garassino 2004). However, it must be reiterated that previous studies on Solnhofen material have only used ultraviolet as a means of fluorescence (Keupp 2007; Schweigert 2011; Tischlinger & Arratia 2013). A distal portion of the left first pereiopod on LB 7 is unseen under UV light but is revealed, albeit faintly, under LSF (Fig. 8). In *Antrimpos speciosus* (LB 8) (Figs 10–11), more detail of the body outline is exposed by LSF under a green 532 nm laser compared with UV and this is emphasised in the magnified portions (Figs 9, 10 B-C, E-F).

As the figures of Solnhofen decapods herein show, the detail revealed under Laser-Stimulated Fluorescence, often surpasses that revealed by ultraviolet fluorescence, with the outline of the entire animal outline being revealed, along with small elements such as antennae and swimmerets (Figs 10, 11). The veneer of the matrix on LB 8 in Figs 10–11 may have obstructed the UV fluorescence and could be prepared further to assist the use of this method. Ultraviolet fluorescence on a specimen of *Alcmonacaris winkleri* Polz, 2008 (LB 9) reveals a faint outline of the animal while recording colour patterning (Fig. 12). Under LSF, green laser light, this colour information is lost but the animal is revealed in its entirety (Fig. 12D). Techniques to rectify this loss of information have been developed using multiple wavelengths (Kaye *et al.* 2015). As Figs 9–12 show, the preserved exoskeleton and the body outline fluoresce at different levels because of the autofluorescent compounds within the arthropod exoskeleton.
Vertebrata

The Solnhofen Limestone has achieved much of its fame as a fossil Konservat-Lagerstätten because of the exceptional preservation of its vertebrate fossils, especially those of volant animals such as the earliest unequivocal fossil bird *Archaeopteryx* and a diverse assemblage of pterosaurs (Beardmore *et al.* 2017; Schwarz *et al.* 2019; Longrich *et al.* 2020) where parts of the flight surfaces are preserved, including wing membranes and feathers (Frey *et al.* 2003; Jäger *et al.* 2018; Benton *et al.* 2019; Kaye *et al.* 2019a; Foth *et al.* 2020; Wilkin 2020). Some of this exceptional preservation can be seen in the examples below (Figs 13–16). The hollow bones of pterosaurs are rarely preserved and through LSF the contrast between preserved bone and the imprints of the skeleton in the matrix is exemplified. The dwarf crocodyliform *Alligatorellus* (Fig. 16) under the blue laser displays soft tissues around the entire body as well as a brighter section at the base of the tail that may represent partial preservation of the caudofemoralis muscle.

However, the exceptional preservation is not restricted to the Tetrapoda, with many fishes also exceptionally well preserved with full articulation and soft part preservation (Konwert 2016; Arratia *et al.* 2019; Ebert & Lane 2019) as well as reptiles (Tennant & Mannion, 2014) (see Fig. 17). Specimens LB 10–13 (Figs 17–20) vary in completeness from articulated individuals to isolated sources of phosphate, resulting in the high level of fluorescence under UV and LSF (Fig. 12).

The specimens studied vary between right and left lateral views, seen in specimens LB 10 and 11 (Figs 17 and 18 respectively). The fluorescence of these specimens (LB 10–13), results from the vertebrate endoskeleton containing high levels of phosphorus (Tischlinger & Arratia 2013). The fossils in white light contrast little with the surrounding sediment as in many other Solnhofen fossils. Indeed, it may be difficult to see some fossils, especially when weathered, without using fluorescence.

Many Solnhofen fossil fish possess a fully ossified skull, fins, neural and haemal spines made apparent through fluorescence along with thin autocentra (Arratia & Schultze 2013). The autocentra surrounding the dark chordocentra are thin and almost translucent and fluoresce under both UV and LSF. Species of the teleost family Orthogonikleithridae are the most common vertebrates in the Solnhofen Limestones (Konwert 2016). The ossification patterns observed on a specimen of *Orthogonikleithrus hoelli* Arratia, 1997 (LB 12) (Fig. 19), correspond with previous studies using UV for fluorescence (Tischlinger & Arratia 2013).

Fish fluoresce well under both UV and LSF and although the difference is often minimal, clearly in Fig. 18 LSF surpasses the UV photograph with an increased fluorescence of both the skull and vertebral column completing the structures seen and this results from the higher flux and subsurface illumination.

**Discussion**

Laser-Stimulated Fluorescence has various applications as illustrated by the numerous examples figured by Kaye *et al.* (2015) including an automated microvertebrate sorting machine developed to reduce time
spent on manual sorting along with demonstrating the potential of LSF to reveal subsurface fossils (Kaye et al. 2015, Figs 10 and 8 respectively). Recently, it has even been applied as part of an automated drone system to seek out fossils on the ground at night (Kaye & Pittman 2020). The technique has revealed geochemical halos around the lost calamus of ‘Archaeopteryx’ (Kaye et al. 2019b), akin to the effect observed in Figs 10–12 where the fluorescence of decapods extends beyond the preserved exoskeleton. The methodology is simple and provides a rapid means for producing high-quality images of fluorescing fossils, allowing for a camera image to be available within 30 seconds of exposure. This exposure time is longer than UV or white light photography as the fluorescence occurs through a fine laser beam scanning an entire specimen rather than the instant illumination under an LED or UV lamp. The same wavelengths of 532nm have been employed previously (Kaye et al. 2015), although this green light is often substituted for blue or violet wavelengths (T. Kaye, pers. comm. 2019). Blue/violet lasers increase the contrast further and reveal fluorescence colour differences akin to those under UV (Schwarz et al. 2019). The 532nm laser in this study results in varying levels of orange fluorescence through the O56 blocking filter that can cause difficulties in distinguishing between different parts of a specimen. As reported by several studies (Mayr et al. 2016; Vinther et al. 2016; Wang et al. 2017; Saitta et al. 2018, Falk et al. 2019; Yang et al. 2019; Pittman & Xu 2020; Serrano et al. 2020), LSF continues to reveal new and exciting details on well-preserved fossil assemblages (Konservat-Lagerstätten) like the Jehol Lagerstätte of northeastern China and the Las Hoyas Lagerstätte of Spain. The high fluorescence we observed in decapods (Figs 8–12) likely results from the autofluorescence present in crustacean exoskeletons (Charbonnier et al. 2017). The Solnhofen specimens analysed here fluoresce brightly due to high levels of phosphate in both original and diagenetic minerals (Wilby et al. 1996). The principal findings in the Solnhofen specimens are that LSF reveals morphology not visible under other UV techniques and more often to a greater degree of clarity and depth.

**Costs**

With reductions to the cost of laser systems, the method could be replicated using a 532nm laser, an LCD power supply and a Zecti 31.5in”/80cm camera slider. This study using an 85mW laser shows the technique can be used with less powerful laser equipment than the 300–500mW laser used by Falk et al. (2016) and Wang et al. (2017). With two wavelengths in tandem, different structures fluoresce, allowing for a more complete picture (Wang et al. 2017), providing an option for further research. The use of a less powerful laser also allows for LSF to be more accessible on the grounds of cost, but with a necessary trade-off in fluorescence signal.

**Conclusions**

The list of non-destructive techniques available to palaeontologists is increasing. X-rays were first implemented in 1896 on fossils from another German Lagerstätte, the Devonian Hunsrück Slate (Hohenstein 2004), and was widely implemented from the 1930s by Lehmann on this site along with other fossil-rich areas like Messel pit and the Solnhofen region. The use of UV has become standard in
many palaeontological studies especially those investigating soft tissue preservation (Hone et al. 2010; Kellner et al. 2010; Cuesta et al. 2015; Schwarz et al. 2019; Hoffman et al. 2020) since its first use in 1926 on fossil vertebrates within the Solnhofen plattenkalks (Simpson 1926). Composite imaging and 3D computer modelling have also been used on Solnhofen fossils and provide a valuable alternative, especially on small and delicate specimens (Haug et al. 2008). Synchrotron Rapid Scanning X-ray Fluorescence (SRS-XRF) combined with chemical images allowed for the mapping of plumage patterns in the iconic Solnhofen bird *Archaeopteryx* (Manning et al. 2013).

LSF was added to the literature as recently as 2015. The use of lower-powered equipment in this study illustrates that the technique can still be employed to good effect and could provide a valuable teaching resource. The method can be performed in as little as 30 seconds, allowing for almost instant data collection. Previous publications have focused entirely on vertebrate material. This study shows the effect of LSF on invertebrates for the first time. In addition, it adds the Solnhofen plattenkalks to the list of Konservat-Lagerstätten where the effects under LSF are now known. This study underscores LSF as an alternative tool to UV for non-destructive palaeontological investigation using extensive comparative figures that show unseen details to an equal and often greater extent than UV.

**Declarations**

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Figure 1

A facies map of the Solnhofen Limestones in Bavaria, southern Germany with the location X indicates the source of the fossils used in this study. Modified from Ebert et al. (2015).
Figure 2

A simplified diagram of the camera and specimen table setup used to perform LSF imaging. 1, The image area of the DSLR camera; 2, Carriage of the camera track holding the laser module; 3, The imaged specimen; 4, Laser illumination using a line generator; 5, 85mW 532nm laser module; 6, DSLR 5300; 7, Tripod for long exposure photography. Not to scale.
Figure 3

Isolated specimen of Lamellaptychus (LB 1), a genus for ammonite mouthparts not associated with a specific taxon, under white light (A-B), UV (C) and LSF (D). A. Direct white light image differentiating between the brown thick spongy layer and the thin crenulated layer beneath; B, oblique white light casting a shadow on the specimen revealing the full outline; C, ultraviolet light of 365 nm wavelength used to display colour differences between the two layers; D, Increased contrast under LSF using a 532 nm green laser allows the full outline of the original shape to be observed. Scale bar = 10mm.
Figure 4

The oppeliid ammonite Neochetoceras bous Oppel, 1863 with a calcified phragmacone, full body chamber outline and preserved aptychi (LB 2). A, oblique white light displaying remnants of the body chamber at the left inside edge; B, UV image displaying the siphuncle, phragmacone and jaw apparatus with colour differences along with possible stomach contents; C, LSF image of Neochetoceras bous
under a 532 nm green laser with phosphatised siphuncle along with a clear boundary between the phragmacone and body chamber unseen in other methods. Scale bar = 10 mm.

Figure 5

The phragmacone of the oppeliid ammonite Fontannesiella prolithographica (Fontannes) (LB 3) with phosphatised siphuncle under white light (A-B), UV (C), and LSF under 532 nm green laser light (D). Note the colour variation seen under 365 nm UV is not consistent with Fig. 6 and may be because of differential preservation. Scale bar = 10 mm.
Figure 6

The oppeliid ammonite Neochetoceras sp. with some phragmacone fluorescence and the ammonite coprolite Lumbricaria attributed to LB 4. A, direct white light recording a slight colour difference around the phragmacone and very little evidence of the coprolite; B, oblique white light; C, UV light fluorescing the coprolite, siphuncle and phragmacone; D, LSF displaying full fluorescence of the phragmacone, increasing the observed contrast. Note the brighter siphuncle under 532 nm green laser light in D, allowing for a greater distinction than under UV light seen in C. Scale bar = 10mm.
Figure 7

The plesioteuthid squid Plesioteuthis prisca with a fluorescent gladius (LB 5). This squid can be seen under oblique (A) and direct white light (B). 365nm UV light fluoresces the rachis in this specimen, showing that the central rachis is raised (C). LSF using the 532 nm laser fluoresces this gladius at different levels with the central vane picked out through its higher fluorescence. Scale bar = 10mm.

Figure 8

An isolated appendage of the fossil prawn Aeger tipularius under different lighting conditions (A-D) (LB 6). A, oblique white light image, casting shadows on the slight relief present; B, direct white light; C, Same specimen under 365 nm ultraviolet fluorescence, highlighting the specimen from the background and recovering some unseen setae that appear broken; D, Laser-stimulated fluorescence image of the
specimen under 532 nm green laser displaying an improvement over the UV image seen in C with the complete leg revealed along with none of the gaps present under UV. Scale bar = 10mm.

**Figure 9**

The eryonid crustacean Cycleryon propinquus von Schlotheim, 1822 under white light (A), 365 nm UV (B) and LSF (C) revealing the full outline of the animal under a 532 nm green laser (LB 7). D and E represent comparisons of the fluorescent techniques on the first pereopod (P1). Note the missing section under UV light is revealed through the subsurface illumination of LSF. Scale represents 1cm. D and E magnified x1.5.
Figure 10

Complete fossil of the penaeid shrimp Antrimpos speciosus Münster, 1839 (LB 8) under different lighting conditions; A, oblique white light showing the split rock revealing the original exoskeleton; B, direct white light; C, UV fluorescence increases the contrast with the background by illuminating the specimen and leaving the matrix dark; D, LSF reveals the entire body outline of the animal and this not seen in the white light or UV photographs. Note the antennulae (atl), swimmerets (sw), walking legs (wl) and urostyle (ur)
which were not fully visible in white light. Abbreviations following C. Haug et al. (2009). The green laser wavelength used on this specimen was 532 nm. Scale bar = 10 mm.

**Figure 11**

Isolated UV (A-C) and LSF (D-F) images from Fig. 10 highlighting the differences in revealed detail. B and E are UV and LSF images of the revealed antennae with clear segmentation visible under LSF. C and F highlight the swimmerets that are fully revealed under LSF with full segmentation. Abbreviations as
above with the addition of the antennular peduncle (ped) from Audo and Charbonnier (2012). Scale bar = 10mm. B and E magnified x2 and C and F magnified x3.6.

**Figure 12**

Specimen LB 9 of the caridean shrimp Alcmonacaris winkleri under oblique white light (A), direct white light (B), UV fluorescence (C) and LSF (D). Notice the faint outline under UV is enhanced through LSF to reveal the full animal, allowing the fossil to be labelled. Scale = 10mm. The features revealed use the same abbreviations as Fig. 9 with the addition of the carapace (ca). Abbreviations following C. Haug et al. (2009). The LSF wavelength used was 532 nm.
Figure 13

Specimen SNSB-BSPG 1935 I 24 of the pterodactyloid pterosaur Ctenochasma gracile under white light (A) and LSF (B). This individual is fully articulated with a blue fluorescent soft tissue body outline and blue cartilage between the light orange bones. A 405 nm blue laser was used to fluoresce this specimen.
Figure 14

Specimen SNSB-BSPG AS I 745a of the pterodactyloid pterosaur Germanodactylus rhamphastinus under white light (A) and LSF (B). The skull and upper torso have an increased contrast under LSF through a 405 nm blue laser, separating the specimen from the surrounding matrix. Specimen badge measures 5cm.
Figure 15

Specimen SNSB-BSPG 1977 XIX 1 of the pterodactyloid pterosaur Germanodactylus rhamphastinus on a counterplate slab under white light (A) and LSF (B). The counterplate slab has some missing bone material where it remains dark under LSF. A 405 nm blue laser was used to carry out LSF and fluoresces any preserved bone material.
Specimen SNSB-BSPG 1937 I 27 of the atoposaurid crocodyliform Alligatorellus beaumonti bavaricus under white light (A) and LSF (B). The colour patterning makes clear distinction possibly between the skeleton and osteoderms and the surrounding soft tissues. Note the brightly fluorescing soft tissues at the base of the tail, possibly phosphatised remnants of muscle tissue. LSF was carried out using a 405 nm blue laser. Scale represents 2cm.
Figure 17

A specimen of the teleost fish Leptolepides sprattiformes Blainville, 1818 (LB 10), under white light (A), UV fluorescence (B) and LSF (C). Not the completed soft tissue outline that is brighter than the ossified material. Gut contents fluoresce brightly at the centre of the body cavity along with a coprolite illuminated at the posterior. The laser used on this specimen was a 532 nm green laser. Scale = 10 mm.
Figure 18

A near complete specimen of the salmoniform fish Orthogonikleithrus hoelli, lacking the dorsal section of caudal fin (LB 11). A, oblique white light photograph showing a complete vertebral column and faint skull and caudal fin elements; B, direct white light image; C, UV fluorescence highlighting the bones of the skull along with the pectoral fin; C, enhanced fluorescence from B showing the caudal fin is incomplete along with fluorescing the entire skull. LSF was carried out with a 532 nm green laser. Scale bar = 20 mm.
Figure 19

Two small specimens of the salmoniform fish Orthognatholeithrus hoelli on the same slab (LB 12) under different light conditions. A, oblique white light photograph with a dark vertebral column and lighter skull and fins present; B, UV photograph with complete skull and tail revealed. Dorsal and pelvic fins are also revealed through fluorescence with illumination from the left; C, LSF image revealing the entire dorsal and pelvic fins on the right specimen that are faint under UV. Note the erect position of the dorsal and pelvic fins on the left and relaxed position on the right. A 532 nm green laser was used to carry out LSF. Scale bar = 10mm.
Figure 20

Specimen LB 13 of the extinct bulldog fish Allothrissops salmoneus (Blainville, 1818) under white light (A), UV (B) and LSF (C) conditions. The fluorescence around the articulated skeleton reveals loose scales along with gular plates and gut contents. A 532 nm green laser was used to carry out LSF. Scale bar = 10 mm.