

A peer-reviewed version of this preprint was published in PeerJ on 15 December 2015.

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Schoenemann B, Clarkson ENK, Horváth G. 2015. Why did the UV-A-induced photoluminescent blue-green glow in trilobite eyes and exoskeletons not cause problems for trilobites? PeerJ 3:e1492
<https://doi.org/10.7717/peerj.1492>

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The calcitic lenses in the eyes of Palaeozoic trilobites are unique in the animal kingdom, although the use of calcite would have conveyed great advantages for vision in aquatic systems. Calcite lenses are transparent, and due to their high refractive index they would facilitate the focusing of light. In some respects, however, calcite lenses bear evident disadvantages. Birefringence would cause double images at different depths, but this is not a problem for trilobites since the difference in the paths of the ordinary and extraordinary rays is less than the diameter of the receptor cells. Another point, not discussed hitherto, is that calcite fluoresces when illuminated with UV-A. Here we show experimentally that calcite lenses fluoresce, and we discuss why fluorescence does not diminish the optical quality of these lenses and the image formed by them. In the environments in which the trilobites lived, UV-A would not have been a relevant factor, and thus fluorescence would not have disturbed or confused their visual system. We also argue that whatever the reason was that calcite was never again used successfully in the visual systems of aquatic arthropods, it was not fluorescence.

Why the UV-A-induced photoluminescent blue-green glow in trilobite eyes and exoskeletons did not cause problems for trilobites?

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Abstract

The calcitic lenses in the eyes of Palaeozoic trilobites are unique in the animal kingdom, although the use of calcite would have conveyed great advantages for vision in aquatic systems. Calcite lenses are transparent, and due to their high refractive index they would facilitate the focusing of light. In some respects, however, calcite lenses bear evident disadvantages. Birefringence would cause double images at different depths, but this is not a problem for trilobites since the difference in the paths of the ordinary and extraordinary rays is less than the diameter of the receptor cells. Another point, not discussed hitherto, is that calcite fluoresces when illuminated with UV-A. Here we show experimentally that calcite lenses fluoresce, and we discuss why fluorescence does not diminish the optical quality of these lenses and the image formed by them. In the environments in which the trilobites lived, UV-A would not have been a

relevant factor, and thus fluorescence would not have disturbed or confused their visual system. We also argue that whatever the reason was that calcite was never again used successfully in the visual systems of aquatic arthropods, it was not fluorescence.

Introduction

Trilobites were the most prevalent mobile invertebrates of the Palaeozoic seas, as known from their fossilised remains. They were arthropods, equipped with a thick shell and highly differentiated compound eyes from the very beginning of their appearance in the fossil record, some 522 million years ago. Trilobites developed a very special optical system, contrasting with that of all other arthropods. For, uniquely in the animal realm they had compound eyes with lenses of oriented calcite rather than of organic material (Towe 1973, Clarkson, Levi-Setti & Horváth 2006, Lee, Torney & Owen 2007, 2012). The use of calcite brings an evident advantage optically, especially for aquatic organisms. The high refractive index of calcite (~ 590 nm: $n_o=1.640-1.660$, $n_e=1.486$) by contrast with that of chitin, the lens material of most other arthropods ($n=1.46$, rarely up to 1.56 (Land & Nilsson 2012)), increases the difference in refractive indices between the visual system of the arthropod and water ($n=1.334$, seawater), and thus facilitates focusing due to strong refraction. Of special interest has been the visual system of a suborder of trilobites, the Phacopina, because their large lenses (diameters of up to 2 mm and more in e.g. *Drotops megalomanicus* Struve 1990) have an elegant internal substructure, which probably corrected lens aberrations (especially spherical aberration), which would otherwise be produced by the thick lenses that phacopid trilobites possess (Clarkson &

Levi-Setti 1975). Although nothing is usually preserved below the level of the lenses, the first known sublensar sensory structures, at a cellular level, have been described in these trilobites very recently (Schoenemann & Clarkson 2013). This raises questions about the specificity of this unique calcitic system, which persisted successfully for more than 250 million years, but was never reinvented again after trilobites became extinct, despite the high advantage of transparency and a high refractive index which allows efficient focusing even under water.

Calcite is a strongly birefringent mineral, and light passing through it in directions other than parallel with the c-axis splits into two rays; producing double images at different depths. At first sight this may seem to be a problem for trilobite vision. But because the difference of paths in the ordinary and extraordinary ray on their way through the lens is smaller than the separation of common photoreceptor units (being usually larger than the receptor diameter), the double images may be irrelevant (Schoenemann & Clarkson 2011).

Another striking characteristic of the mineral calcite, apart from birefringence, is photoluminescence. The photoluminescence is usually related to impurities of organic material or minerals, such as magnesium, manganese, iron etc. as well as cracks (Machel 1985, Machel et al. 1991, Pedone et al. 1990). Natural calcite fluoresces when it is illuminated with light of certain wavelengths, as for example UV-light, and the colour of this fluorescence depends on the character of the particles the calcite includes. The energy of the incident light is able to excite susceptible electrons within the atomic structure of the mineral. They leave their position and jump to higher orbits of the atomic structure. Falling back they release a small amount of energy visible as light, and producing a kind of 'glow'. The colour of this 'glow' often

is different from the colour of the incident light, and depends on the composition of the calcite, while the 'glow' continues as long as the mineral is illuminated. The colour of the glow depends on the orbit from which the electron returns to its original position. In contrast, during phosphorescence, the light is 'stored' for a while inside the atomic structure, the system becomes 'charged', and releases the energy more slowly than during the fluorescent process. The excited electron also returns to its position inside the atomic structure but it undergoes certain intersystem levels, while its state of spin turns to a higher spin multiplicity, normally a triplet state. These transitions take time in the order of milliseconds, but can also persist in some materials for minutes or even hours. In our probe, the phosphorescence, seen in a biological time scale (milliseconds), disappears as soon as the light vanishes. While calcite shares this property of showing fluorescence with numerous other natural minerals, such as fluorites or opals, and synthetic minerals also (Nakamura et al 2013) at a first glance it seems quite extraordinary to find a presumably fluorescent mineral element in the morphology of a biological system, especially a visual system.

Calcium carbonate exists in many biological systems. For example, in the form of calcite it is reported from light-sensitive systems in brittle stars (Aizenberg et al 2001), the shells of brachiopods, ostracodes and other crustaceans (Xia et al. 1997). On the other hand, the shells of many kinds of molluscs are built of aragonite, a form of calcium carbonate with a crystal lattice different from that of calcite, and typical for the exoskeletons of corals, and some serpulids. Calcium carbonate (calcite) is not known so far in image-forming structures, except in the trilobites.

85

86 Bioluminescence occurs widely in living systems, especially in marine vertebrates,
87 invertebrates, some fungi, and many microorganisms, but not in land vertebrates and higher
88 plants. There is here a distinction between primary bioluminescence, where the organism itself
89 generates the light, and secondary bioluminescence, where the light is produced by symbiotic
90 microorganisms, which are, of course themselves, primary bioluminescents. A very common,
91 basic system is the oxidation of luciferin by the enzyme luciferase, there are other enzymes
92 involved such as superphotoxidase in fungi (Shimonura 1992, Desjardin et al. 2008) is involved,
93 or aequorin in the jellyfish *Aequorea victoria* (Hastings 1983, Kendall & Badminton 1998,
94 Shimonura 2005, Gruber & Pieribone 2007, Meyer-Rochow 2009, Haddock et al. 2010, Sparks et
95 al 2014). Bioluminescence is used to attract mating partners, for defence, warning, mimicry,
96 and illumination or as counterillumination balancing the residual downwelling light to cloak the
97 silhouette from upward-looking predators, as was recently reported for bioluminescent sharks
98 (Claes et al. 2014). Whether bioluminescence is useful, especially fluorescence in a visual organ,
99 such as is caused by UV-light in the calcitic lenses of the dioptric apparatus in trilobite
100 compound eyes, may be worthwhile to consider further.

101 The precise analysis of different trilobite lenses has shown that during diagenesis the
102 composition of the calcitic lenses of different trilobites has been altered (Lee,Torney & Owen
103 2007, 2012), as it becomes very evident in the meanwhile famous red trilobites with green
104 eyes, from Morocco, which had undergone a silicified preservation rather than a fossilisation in
105 limestone as is more or less usual (Klug, Schulz & De Baets 2009). The Hunsrück Slate is well-
106 known for its exceptional preservation and that calcium carbonate is often dissolved or

replaced. As is shown here, however, we still see fluorescence even today, so it seems allowed to assume that there exists no pervasive diagenetic influence on this system. It will not be possible, however, to reconstruct the precise original mineral composition of the lenses. Consequently, the actual character of the fluorescence in the lenses of trilobite compound eyes during the life-times of the trilobites remains unknown, but some discussion of the relevance of the potential phenomenon of fluorescence in these ancient calcitic lenses, in principle, would seem desirable. Actually, there are three main questions which it seems worthwhile to answer:

1. Do the lenses of trilobite eyes, after all this theoretical discussion, really show fluorescence?
2. What are the optical and sensory consequences of fluorescence, if this is indeed what they actually show?
3. Is the reason, why calcite has not been used more often in aquatic optical systems, the fact that it is fluorescent?

MATERIALS AND METHODS

Because most trilobite exoskeletons fossilised in limestones are largely composed of calcite, experiments for investigating the photoluminescence of calcite lenses were performed on a species which normally fossilises in a somewhat different way. The specimens used here come from the Bundenbachschiefer of the Lower Devonian of the Hunsrück region, Germany. In these trilobites, the sulfur released from proteins, together with iron from the ancient mud formed pyrite, while the lenses of pure calcite stayed as they were. The phacopid trilobite

Chotecops ferdinandi (Kayser, 1880) (Fig. 1a) is very abundant at this location and possesses large (~7mm) compound eyes. Lens preservation, however, is extremely rare, because the lenses normally fall out of the fossil, and cavities remain where the lenses had been. Even so, very occasional examples are found such as the two isolated eyes of moulted specimens used here, each showing the phenomenon independently (Fig. 1c-f). Detailed reports about the age and setting of the Hunsrück Slate fauna, taphonomy and lithostratigraphy are given in e.g. Schindler et al. (2002), Kühl et al. (2012) and De Baets et al. (2013). The specimens are housed in the collection of the Geological Institute of the University of Cologne (now Institute of Geology and Mineralogy). The museum numbers are GIK 2118 and GIK 2119. They were illuminated with a peak-wavelength of ~365nm (UV-A: specification of the manufacturer) from a source of low energy (6V, 4W, 40mA, ETT Comp. Braunschweig, Germany, specification of the manufacturer) and photographed (Panasonic DMC-TZ10). The width of the spectrum of the light source is unknown and is not relevant for showing the principal phenomenon of fluorescence in the calcitic lenses of phacopid trilobite compound eyes.

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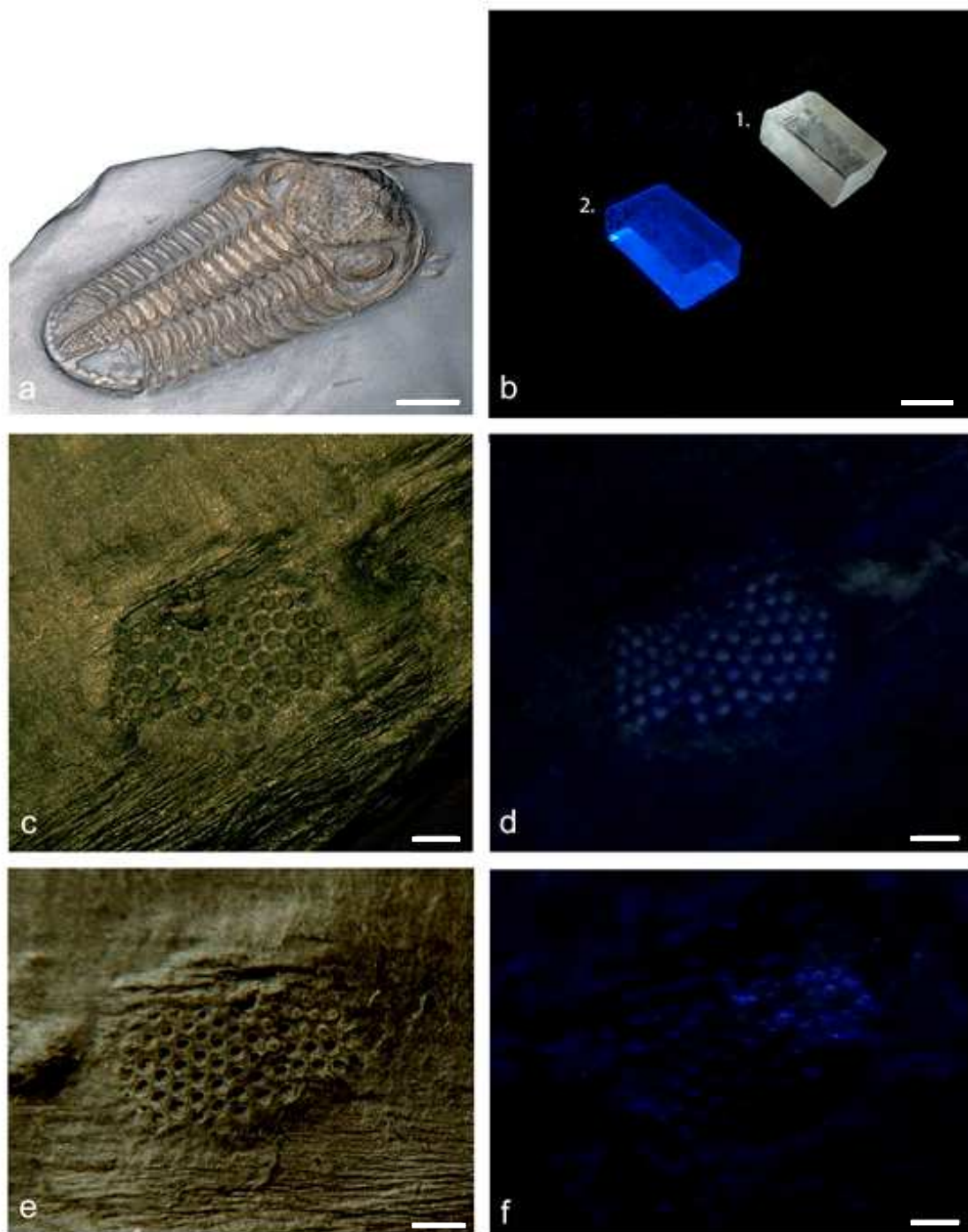
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155 **RESULTS**



156

157 **Figure 1.** The glow in the calcitic lenses of a phacopid trilobite's eye.

158 (a) *Chotecops ferdinandi* (Kayser, 1880), Bundenbachschiefer, Lower Devonian, Location:
159 Grube Eschenbach, Hunsrück, Germany, scale bar ~1cm (housed in the collection of Steinmann
160 Institute, University of Bonn still open, curator on field work). (b) 1. Calcite crystal (~3cm), scale
161 bar ~1cm. 2. fluorescent when illuminated with ~365nm under water. (c) Isolated moult of a
162 *Chotecops* compound eye with lenses preserved (GIK 2118). (d) The same showing fluorescence
163 in the calcitic lenses of the trilobite compound eye when illuminated with UVA-light (~365nm).
164 (e) Isolated moult of a *Chotecops* compound eye with lenses preserved (GIK 2119). (d) The same
165 showing fluorescence in the calcitic lenses of the trilobite compound eye when illuminated with
166 UVA-light (365nm). (b-f) scale bar ~1mm.

167

168 When illuminated with UV-A light (365nm) the remains of the calcitic lenses glow with a blue-
169 greenish light as long as they are illuminated, while other parts of the eye, which are not of lens
170 material, remain (more or less) dark. Both of the extremely rare specimens show the
171 phenomenon in the same way and independently.

172

173 Discussion

174 The fact that the material of trilobite lenses was primary calcite, as proposed by Towe (1973),
175 has been unequivocally confirmed, the lenses of all known species were originally calcitic,
176 independently of how they have been preserved (Clarkson 1975, 1979, 1997, Clarkson et al.
177 2006). This understanding has been strengthened by the use of mineralogical methods and

particularly by the use of Electron Backscattered Diffraction (EBSD) technology (Lee et al. 2007, 2012). These Lower Devonian compound eyes investigated here are almost 400 million years old. As already mentioned, the mineral content may have changed during preservation and possible recrystallisation, and consequently the colour of fluorescence and its intensity may have changed. Whereas it will probably never be possible to reconstruct the original composition precisely, the potential to generate the phenomenon itself in principle, however, is clearly shown in Figure 1, where the calcitic lenses so evidently fluoresce. So the first and basic question, whether there is really some potential in the calcitic lenses of trilobite eyes to produce fluorescence, can be answered positively.

2. What are the optical and sensory consequences of such fluorescence?

It seems necessary to consider firstly what would be the consequences for the visual system, if we had a pure perception of the UV-A light, and no other.

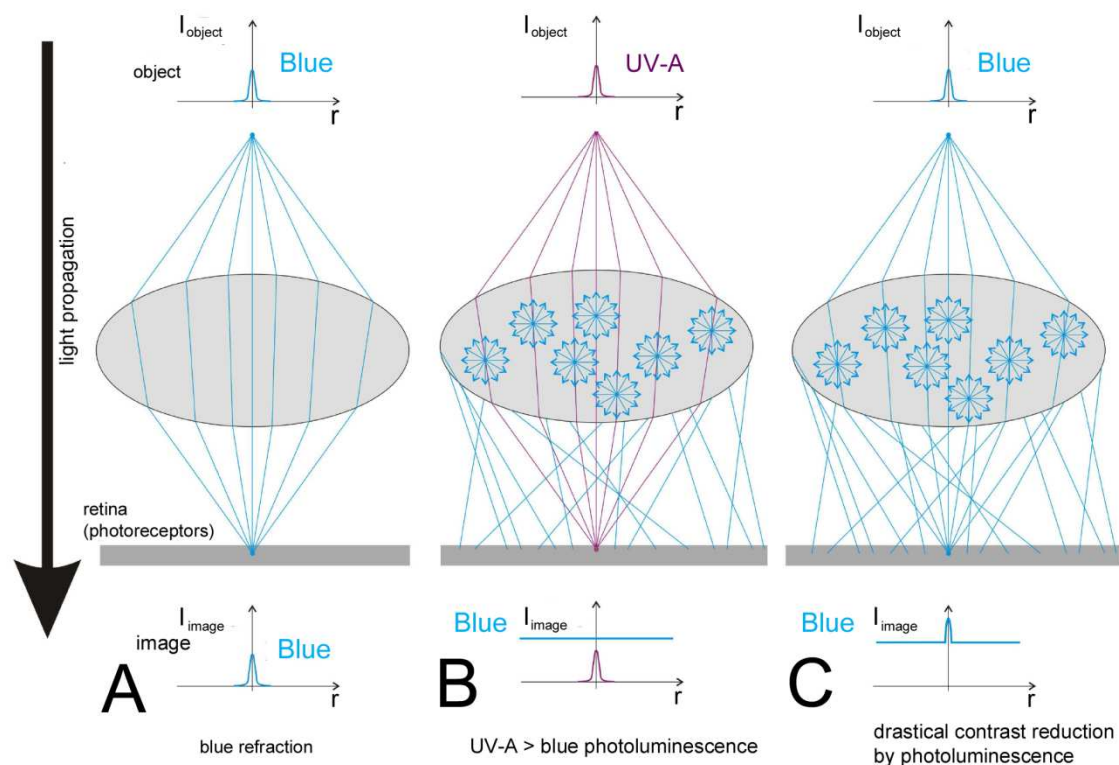


Figure 2 The optical problem caused by the UV-A-induced photoluminescent diffuse blue light in the image formation by a dioptric apparatus (for explanation see text).

As figure 2a shows, the normal function of a lens is to focus incident light to one point. We do not know exactly, what the underlying sensory system in a trilobite's compound eye actually was. It is rather probable, that under each lens of the compound eye, which from outside is recognisable as a facet, was a so-called ommatidium, as we find it in apposition compound eyes of many diurnal arthropods living today, such as dragonflies or bees. It is the oldest system of compound eyes; more advanced systems adapted to dimmer light conditions probably did not

evolve before the Devonian (Gaten 1998). In the apposition eyes the light is focused through a normally chitinous lens, or structure functioning as such, onto a central light guiding structure, the so-called rhabdom, which is part of several (often eight) photoreceptor cells. In the rhabdom lie the photopigments, and the energy of the incident light alters the sterical form of the photopigments to evoke an electrical signal, which can be processed by the nervous system of the organism. The ommatidia are isolated from each other by pigment cells. Because the rhabdom integrates all optical inputs inside the angle of view of the ommatidium, there results over the entire compound eye a mosaic-like image. The higher the number of facets, the more acute is the image, in the same way that pixels contribute to a computer graphic, and the smaller the field of view of each ommatidium actually is. An indication that trilobites had a kind of apposition compound eye was described recently using x-ray tomography and synchrotron radiation in phacopid trilobites (Schoenemann & Clarkson 2013). An alternative to this system is the establishment of a small retina, a layer of receptors below the lens, as we know it among arthropods from myriapods and many chelicerates. If there was a third alternative, it would not yet be known.

In principle, this mosaic-like character of the image formed by an apposition eye should more or less be retained by any fluorescent pattern of the compound eyes' lenses generated by an inhomogenous UV-light distribution in the environment, but because in sum all points of fluorescence inside the lens cause a high loss of contrast, this principle cannot be adopted entirely, as we shall see.

Figure 2 shows what happens, when UV-A light enters the calcitic lenses of a trilobite. An object point is characterized by the intensity function $I_{object}(r)$, where r is the radius measured perpendicularly from the optical axis. It should be projected onto the light-sensitive receptor plane as a sharp image point, the function $I_{image}(r)$ of which is similar to $I_{object}(r)$. Sharpness means that the narrow and high intensity peak of $I_{object}(r)$ is transferred by the dioptric apparatus as a similarly narrow and similarly high intensity peak of $I_{image}(r)$ as seen in Fig. 2A for blue light, characteristic of the semi-monochromatic optical environment of trilobites.

A point source of UV-A light is similarly imaged onto the retina as shown in Fig. 2B. But UV-A induces blue(-green) light in the bulk calcite medium of the lens. This UV-A-induced blue light propagates in all possible directions from its numerous point sources in the lens. After refraction on the lens surface, this diffuse blue light reaches the sensory system below, where it forms a relatively intense, practically homogeneous blue background light field $I_{blue}(r) =$ constant (Fig. 2B). Thus, the sharp-peaked object point with $I_{object}(r)$ is projected as a wide blue circular spot with a small intensity peak in its center, as shown in Figure 2C.

This would happen to each of the tesserae in the mosaic-like vision of a trilobite compound eye with an assumed apposition eye system. It would destroy the integrating properties of the rhabdom because of a loss of intensity in its signal received. Over the whole compound eye this would result in a loss of contrast.

If we had a retinal system below the lens, this mechanism would help to supply all receptor cells of this visual unit with light – but then the question rises as to why there is a, sometimes probably even sophisticated lens with a central focusing (Clarkson & Levi-Setti 1975). A light distribution as results to $I_{image}(r)$ would be of help just in a combined system, something with a

244 centralised visual system like a fovea/or ommatidium and peripherally a supportive system with
245 retinal receptor cells.

246

247 Another disadvantage might be that the UV-A light from nearly all angles would enter the lens,
248 and thus the fluorescence would be roughly equal in all lenses of the eye, irrespective of which
249 part of the visual field the light came from. Any image formation would be corrupted, the
250 details of the environment would be no more resolved than just light or no light.

251

252

253 It is well known, that clear seawater has a transmission maximum at about 470nm (Figure 3A),
254 so everything a trilobite living at a depth deeper than a few meters saw, would appear in a blue
255 greenish light. UV-A light is attenuated roughly three times faster than blue light, making
256 underwater environments contain much less UV than terrestrial habitats. Already above the
257 water surface there is much less UV than blue – on a sunny day there is about four times as
258 much blue (470nm) as there is UV (365nm).

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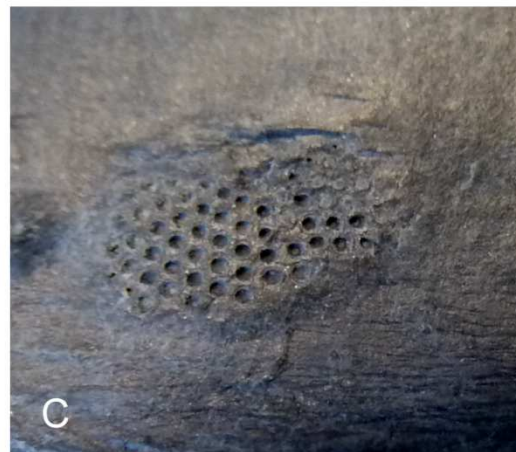
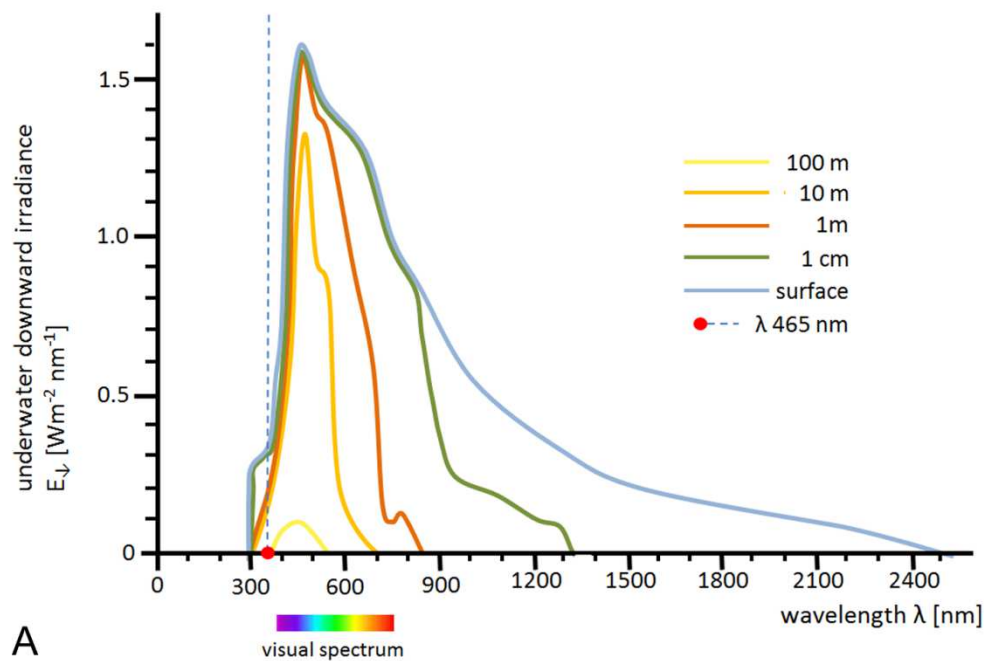


Figure 3 Underwater downward irradiance and fluorescence of trilobite lenses under UV-A light and day-light conditions.

(a) Underwater downward irradiance (changed and simplified after Wozniak & Dera 2007). (b) Isolated moult of a *Chotecops* compound eye with lenses preserved (GIK 2118) showing a very

266 slight fluorescence in the calcitic lenses of the trilobite compound eye when illuminated with
267 UV-A light (~365nm) under day-light conditions. (c) Isolated moult of a *Chotecops* compound
268 eye with lenses preserved (GIK 2119) showing a very slight fluorescence in the calcitic lenses of
269 the trilobite compound eye when illuminated with UV-A light (365nm) under day-light
270 conditions, but not as evident as in b. (b, c) Scale bar ~1mm.

271 The photoluminescence of the calcitic lenses in trilobites, may have enhanced the width of the
272 exploitable spectrum of vision of their bearers, transforming the UV-light to a fluorescence. We
273 do not know the exact contents of impurities of the original calcitic lenses, thus nothing about
274 the likely exact colour of a potential fluorescence. By physical reasons mentioned before it may
275 be assumed that the early photoreceptors were sensitive to blue light, as are most
276 photoreceptors of aquatic animals still today, which would match a blueish green fluorescence
277 as shown in our experiment. If, at this early time in the evolution of complex marine animals,
278 specialised UV-receptors had not yet originated, by transforming UV-A by fluorescence of the
279 lenses overlying the receptor system into blue-greenish light, it might have been possible to
280 'catch' these shorter wavelengths. This would extend the normal range of wavelengths
281 available for vision, but without requiring specialised blue-green receptors. An argument
282 against this facility is that there seems to be a general rule in (underwater) visual ecology, that
283 where UV-A is available in a given optical environment, there the animals have also UV-A
284 sensitive photoreceptors, and only those animals do not have UV-A receptors which live in a
285 UV-A deficient environment. So, why should trilobites would be an exception of this rule, and
286 all the more, since during their 270-million-year history they could have been able to develop
287 UV-A sensitive receptors, similarly to those of many recent marine animals. Furthermore,

probably it would have 'cost' less to establish UV-A sensitive cells rather than a calcitic lens. But although we do have the calcitic lenses, we know nothing about the properties of the receptor cells below, and it is likely that any UV-A-induced fluorescence in this system would have produced images of very poor quality because of a drastically reduced contrast.

Regarding other visual systems of today, it is well known that in all known recent visual systems the amount of light scattered diffusely in the dioptric media (cornea, lens, crystalline cone, etc.) is minimized. In the human eye, for example, light is scattered diffusely in the vitreous body, which gives a non-imaging interior light field, greatly disadvantageous for image formation. One of the functions of the retinal pigment epithelium (containing melanin between the chorioid and retina) is to absorb this vitreous-scattered light.

The cuticular microstructure of the trilobites' exoskeleton has been explored by several workers. It is generally agreed that the cuticle consists of the following layers (i) a very thin, originally organic layer, not often preserved, and sometimes phosphatised, (ii) a thin outer layer, often prismatic, with the crystallites arranged perpendicular to the surface. This outer layer, in *Asaphus* is about 1/15th of the total thickness of the cuticle (Dalingwater 1973) (iii) a much thicker principal layer with distinct laminations, parallel with the outer and inner surfaces. This, like the outer layer, consists of low magnesian calcite (Wilmot & Fallick 1989). Dalingwater & Miller (1977) note that in the principal layer "Individual calcite crystals are difficult to resolve, but roughly shaped perpendicular plates of calcite [...*normal to the cuticle surface* ...] are prominent... in some cases pierced by canal-like elements". Likewise Dalingwater

et al. (1991) comment that the principal layer "consists of fine crystallites, presumably of calcite, sometimes with their long axes arranged roughly perpendicular to the cuticle surface". Wilmot (1990) notes that trilobite cuticles were able to resist both tensile and compressive forces. The outer, prismatic layer was able to resist compressive forces acting normal to the surface. The principal layer, on the other hand, with its small crystals, acted as a crack-stopper, as well as giving bulk to the exoskeleton.

So, in other words, the principal layer consists of small calcite crystals, sometimes with a rough orientation perpendicular to the surface.

'Calcite in trilobite eyes was likewise orientated so that its c-axis was parallel to the optical axis of the lens and perpendicular to the surface. This ordered calcite orientation minimized the optical problem caused by the birefringence of calcite. The eyes are only a specialised part of the exoskeleton, and the orientation of the c-axes in the lenses are concordant with the overall structure of the cuticle. However, the calcite crystals in the exoskeleton should also transfer UV-A to blue-green light. Thus, the whole body surface of trilobites illuminated by UV-A light should emit faint blue-green light, which could be very disadvantageous due to camouflage disruption: a trilobite emitting blue-green light would be visually very striking both for their prey and predators. Unfortunately the emission of this blue-green light in our fossils is so low that it cannot be photographed.

Thus, if calcite in lenses as found in the optical apparatus of trilobite compound eyes had such disadvantageous properties for any visual quality, and even the exoskeleton under UV-light may have been somewhat luminescent, evolution should somehow have eliminated these disruptive phenomena. A simple method to improve the quality of trilobite vision would have been to avoid the use of calcite in the dioptric apparatus altogether – Recent arthropods use chitin instead.

There thus remain interesting questions to be answered. Was fluorescence the reason why calcite has not been used more often in aquatic optical systems? And: Why the UV-A-induced photoluminescent blue-green glow in trilobite eyes and exoskeletons did not cause problems for the trilobites?

A first strategy to escape from fluorescence would be to produce a calcite so pure that it does not contain any impurities. Whether, however, this was possible for a biological system remains doubtful.

Another effective strategy would be to avoid the UV-A light itself. Many trilobites probably have lived a crepuscular or nocturnal life (Clarkson 1998), when their light environment was UV-A deficient. In particular the early trilobites of the Cambrian and probably their predecessors were bottom dwellers. The invasion of the pelagic and planktonic realm by trilobites did not begin before the Furongian (upper Cambrian) and only was truly under way in the early

Ordovician and later (McCormick & Fortey 1998, Tortello & Esteban 2003, Schoenemann et al. 2010, Tanaka et al. 2015), during the Great Ordovician Biodiversity Event.

Figure 3a shows the well-known optical fact that both UV-A/B/C and infrared light are strongly absorbed by (sea)water. As light propagates deeper and deeper into water, both the short (UV) and long (IR, red, green) wavelengths are quickly absorbed, and depending on the water type, after a few decimeters/meters only quasi-monochromatic blue (~475 nm) light remains. Due to this strong wavelength-selective absorption of water, the UV-A intensity of light is practically zero in water deeper than a few m or dm. The majority of trilobites surely lived deeper in the sea than a few m/dm.

Furthermore, many of the early trilobites presumably lived on organic material on or in the sea-floor sediment, and many of them preferred muddy ecosystems. When the mud was perturbed the water would become turbid. The optical haziness in sea water is caused by fine particles which scatter and absorb UV-A light very strongly. The intensity of the scattered light depends on the fourth power of the frequency, so blue and UV-light are scattered much more strongly than red light.

In consequence, the photoluminescence of their calcite lenses was visually irrelevant, because it was (more or less) not present in the early trilobites' environment.

Finally one should bear in mind the conditions of radiance during the Palaeozoic, when the trilobites were living. It is well known that due to the ozone layer being deficient or absent during the Archean, high energy radiation was able to penetrate more deeply into oceans than

it does at present, and thus the potential damage rates to DNA were magnitudes higher than today. DNA-damage must have been the principal factor for UV-induced mortality in the Archean oceans (Cockell 1998, 2000 a,b, Cockell & Horneck 2001). Thus at 5m depth the potential DNA-damage rate may have been 2 orders of magnitude higher than today, and still one order higher at 15m depth (Cockell 2000a). A quite rapid change started probably ~800 million years ago [Ma] (Qiu 2014), and by at least 700 Ma oxygen levels might have been sufficient for respiration in metazoans (Margulis, Walker & Ramblerer 1976, Bekker et al. 2004, Hessen 2008). Having just about achieved an almost modern atmosphere ~520 Ma, and probably due to the availability of certain minerals for the construction of shells of modern type (Cook & Shergold 1984), the 'Cambrian explosion' became possible, and it was during this time that most modern clades originated (Margulis, Walker & Ramblerer 1976, Cowen 2005, Marshall 2006, Hessen 2008, Erwin et al. 2011). Trilobites appear in the Lower Cambrian among the oldest arthropod fossils, well equipped with a hard shell and complex compound eyes. As for many organisms of this era, the origin of trilobites probably lies before the 'Cambrian explosion' further back in the Proterozoic, though without any fossil record, and we know little of the circumstances of radiation during the early evolution of the compound eyes of trilobites and their predecessors. Whether the invasion of the UV-A-deficient ecological niche as described was a consequence of the calcitic lenses, remains open, but is unlikely. It seems more realistic to assume that trilobites tracked regions rich in organic material easily to be digested, such as down in the muddy grounds of the ocean.

While during the late Proterozoic/early Palaeozoic the ozone levels rose, UV-B and UV-C then were shielded almost completely, while UV-A was able to penetrate before this change, as it

still does, and the amount of UV-A is comparable to that of today. But it surely is a good estimation to say, that of the UV-A light just a small part causes fluorescence, while the rest passes through the thin lenses ($\sim 200\mu\text{m}$), while UV-A itself is just a small part of the light incident reaching the receptors. Furthermore, the percentage of non-UV-A light with respect to UV-A/B/C-light, both at present and at the beginning of the Cambrian was high enough, that any ill-effects of fluorescence due to a low amount of UV-A were very minor relative to light of longer wavelengths transmitted through the lens. Figure 1c and 1e show the eyes and the lenses in 'normal' light, where fluorescence does not become apparent or does not occur. A slight blue fluorescence, however, in figure 3b, is evident where the same eyes are illuminated under day light, with UV-A light in the same way as under the same dark conditions of figure 1. One has to notice, of course, that under these conditions decades more of energy influenced the lenses. In the second specimen under the same conditions, fluorescence was not as evident, nor was it possible to cause any trilobite exoskeleton to glow. Whether the calcitic lenses originated even further back in time, when the UV-content was higher, is without any confident fossil record.

In this context there should be mentioned, however, the publication of Frank & Widder (1996). Electrophysiological experiments on several species of deep-sea shrimp revealed unexpectedly high spectral sensitivity to UV light. Subsequent measurements of downward irradiance at 380nm showed that UV of this wavelength was still detectable at 500 to 600 m, and this is indeed the depth at which these crustaceans live. So UV-relevant phenomena seem to occur at deeper depths, but although the energy of the UV-light may be high enough to switch on highly sensitive receptor cells, it is possibly too low to evoke efficient fluorescent signals.

417

418 In summary, it is possible to answer the questions raised at the beginning.

419 The results show that there is a real potential for the lenses of trilobite eyes to show
420 fluorescence (Fig. 1a). The optical and sensory consequences of fluorescence, as we have
421 discussed, would, however, have been disastrous to the quality of vision because of a high loss
422 of contrast (Fig. 2).

423 Fluorescence, however, is not the reason, why calcite has not been used more often in aquatic
424 optical systems. There are several reasons for that. The disadvantages of an optical system
425 under UV-A light easily can be avoided by invading ecological niches which UV-light cannot
426 influence – as indeed the early trilobites did. They were bottom dwellers, living on muddy sea
427 floors where the water could readily become turbid when the substrate was stirred up, as
428 would be expected for trilobites searching for organic material. Under such conditions, light of
429 short wavelength was effectively scattered and absorbed. But the reason to invade this hazy
430 part of the ocean in the first place, to where UV-A/B/C never passed through, was the
431 availability of appropriate nutrients. This also answers the question, why the UV-A-induced
432 photoluminescent blue-green glow in trilobite eyes and the camouflage-breaking properties of
433 their exoskeletons did not cause problems for trilobites – it was not present in the environment
434 that the early representatives preferred. So the calcitic lenses, with their great ability to focus
435 light under water due to their high refractive index, probably originated in conditions where
436 adverse stimulation caused by UV-light fluorescence was not a factor. Thus, the also important
437 question, whether the fluorescent properties of calcitic lenses were a primary reason why

calcite was never again used in underwater visual systems, can be answered very firmly with:
no.

AUTHOR CONTRIBUTIONS B.S. did the experiments and did the calculations, B.S. and E.C. wrote the manuscript. G.H. gave excellent comments in his review and added figure 2 and some very helpful and essential paragraphs to the manuscript, especially clarifying the phenomenons of fluorescence physically and their function in the trilobite's environment.

ACKNOWLEDGEMENTS We thank Wouter Südkamp, Bundenbach, for the specimen, Kai Jäger, Georg Oleschinski and Gabi Kühl (all Steinmann Institute, University of Bonn) for the photograph of the trilobite in figure 1a, and we are greatly indepted to thank one anonymous reviewer, Gabor Hórváth as a second, Clare Torney as a third reviewer and the academic editor Kenneth De Baets for their helpful and constructive comments. After the 1st round of revision, Gábor Hórváth joined us as a co-author due to his extensive input and original additions to the research.

There are no conflicts of interests of the authors.

REFERENCES

1. Aizenberg J., Tkachenko A., Weiner S., Addadi L. and Hendler G. (2001). Calcitic microlenses as part of the photoreceptor system in brittlestars. *Nature* 412, 819 – 822.

2. Bekker A., Holland H.D., Wang P.-L., Rumble, D., Stein H.J., Hannah J.L., Coetzee L.L. and Beukes N.J. (2004). Dating the rise of atmospheric oxygen. *Nature* 427, 117-120.

3. Claes J.M., Nilsson D.-E., Straube N., Collin S.P. and Mallefet J. (2014). Iso-luminance counterillumination drove bioluminescent shark radiation. *Scientific Reports*, 4: 4328.

4. Clarkson E.N.K. (1975). The evolution of the eye in trilobites. *Fossils & Strata* 4, 7-31.

5. Clarkson E.N.K. (1979). The visual system of trilobites. *Palaeontology* 22(1), 1-22.

477 6. Clarkson, E. N. K. (1997). In The eye, morphology, function, and evolution. In *Treatise on*
478 *Invertebrate Palaeontology*. Part O *Trilobites*, revised. Whittington, H. B. et al. (Edts.) (The
479 Geological Society of America and University of Kansas, Boulder Colorado, and Lawrence,
480 Kansas), pp. 114-132.

481

482 7. Clarkson E.N.K. (1998). *Invertebrate Palaeontology and Evolution*. Blackwell, Malden (USA),
483 Oxford (UK), Carlton (Aus).

484

485 8. Clarkson E.N.K. and Levi-Setti R. (1975). Trilobite eyes and the optics of Des Cartes and
486 Huygens. *Nature* 254 (5502), 663-667.

487

488 9. Clarkson E.N.K., Levi-Setti R. and Horváth G. (2006). The eyes of trilobites, the oldest
489 preserved visual system. *Arthropod Structure and Development* 35(4), 247–259.

490

491 10. Cockell C.S. (1998). Biological effects of high ultraviolet radiation on early earth. *Journal of*
492 *Theoretical Biology* 193(4), 717-729.

493

494 11. Cockell C.S. (2000a). The ultraviolet history of the terrestrial planets: Implications for
495 biological evolution. *Planetary and Space Science* 48(2-3), 20-214.

496

497 12. Cockell C.S. (2000b). Ultraviolet radiation and the photobiology of earth's early oceans.

498 *Origins of Life and Evolution of the Biosphere* 30(5), 467-499.

499

500 13. Cockell C.S. and Horneck G. (2001). The history of the UV radiation climate on earth –

501 theoretical and space based observations. *Photochemistry and Photobiology* 73(4), 447-451.

502

503 14. Cook P.J. & Shergold J.H. (1984). Phosphorus, phosphorites, and skeletal evolution at the

504 Precambrian–Cambrian boundary. *Nature*, 308(5956), 231–236.

505

506 15. Cowen R. (2005). *The History of Life*. Blackwell, Malden (USA), Oxford (UK), Carlton (Aus).

507

508 16. Dalingwater J. E. (1973). Trilobite cuticle microstructure and composition. *Palaeontology*

509 16(4), 827-839.

510

511 17. Dalingwater J. E. and Miller J. (1977). The laminae and cuticular organisation of the trilobite

512 *Asaphus raniceps*. *Palaeontology* 20 (1), (1977)

513

514 18. Dalingwater J. E, Hutchinson S. J., Mutvei H. and Siveter, D. J. (1991). Cuticular

515 ultrastructure of the trilobite *Ellipsocephalus polytomus* from the Middle Cambrian of Oeland,
516 Sweden. *Palaeontology* 34(1), 205-217.

517

518 19. De Baets, K., Klug, C., Korn, D., Bartels, C. and Poschmann, M. (2013). Emsian Ammonoidea
519 and the age of the Hunsrück Slate (Rhenish Mountains, Western Germany). *Palaeontographica*
520 A, 299 (1-6), 1-113.

521

522 20. Desjardin D.E., Oliveira A.G. and Stevani C.V. (2008). Fungi bioluminescence revisited.
523 *Photochemical & Photobiological Sciences* 7(2), 170–182.

524

525 21. Erwin D.H., Laflamme M., Tweedt S.M., Sperling E.A., Pisani D., and Peterson K.J. (2011). The
526 Cambrian Conundrum: Early Divergence and Later Ecological Success in the Early History of
527 Animals. *Science* 334,1091-1097.

528

529 22. Frank F. and Widder E.A. (1996). UV light in the deep-sea: *In situ* measurements of
530 downwelling irradiance in relation to the visual threshold sensitivity of UV-sensitive
531 crustaceans. *Marine and Freshwater Behaviour and Physiology*, 27 (2-3), 189-197.

532

533 23. Gaten E. (1998). Eye structure and phylogeny: is there an insight? The evolution of
534 superposition eyes in the Decapoda (Crustacea). *Contributions to Zoology*, 67(4), 223-235.

535

536 24. Gruber D. F. and Pieribone V. (2007). A glow in the Dark: The Revolutionary Science of
537 Biofluorescence. Harvard University Press, pp. 263.

538

539 25. Haddock H.D., Moline M.A. and Case J.F. (2010). Bioluminescence in the Sea. *Annual Review*
540 *of Marine Science* 2, 443-493.

541

542 26. Hastings J.W. (1983) Biological diversity, chemical mechanisms, and the evolutionary origins
543 of bioluminescent systems. *Journal of Molecular Evolution* 19(5), 309–21.

544

545 27. Hessen D.O. (2008). Solar radiation and the evolution of life. In *Solar Radiation and Human*
546 *Health*, E. Bjertness ed. (The Norwegian Academy of Sciences and Letters, Oslo), 123-136.

547

548 28. Kayser E. (1880). Ueber *Dalmanites rhenanus*, eine Art der Hausmannigruppe, und einige
549 andere Trilobiten aus den älteren rheinischen Dachschiefern. *Zeitschrift der Deutschen*
550 *Geologischen Gesellschaft* 32(1), 19-24.

551

552 29. Kendall J.M. and Badminton M.N. (1998). *Aequorea victoria* bioluminescence moves into an
553 *exciting new era. Trends in Biotechnology* 16(5), 216–224.

554

555 30. Klug C., Schulz H. and De Baets K. (2009). Red Devonian trilobites with green eyes from
556 Morocco and the silicification of the trilobite exoskeleton. *Acta Palaeontologica Polonica* 54(1),
557 117-123.

558

559 31. Kühl G., Bartels C., Briggs D.E.G. and Rust J. (2012). *Visions of a Vanished World: The*
560 *Extraordinary Fossils of the Hunsrück Slate*, Yale University Press, New Haven, London.

561

562 32. Land M.F. and Nilsson D.E. (2012). *Animal Eyes*. Oxford University Press, Animal Biology
563 Series, Oxford.

564

565 33. Lee M.R., Torney C. and Owen A.W. (2007). Magnesium-rich intralensar structures in
566 schizochroal trilobite eyes. *Palaeontology* 50(5), 1031-1037.

567

568 34. Lee M., Torney C. and Owen A.W. (2012). Biomineralisation in the Palaeozoic oceans:
569 evidence for simultaneous crystallisation of high and low magnesium calcite by phacopine
570 trilobites. *Chemical Geology* 314(17), 33-44.

571

35. Machel H.-G. (1985). Cathodoluminescence in calcite and dolomite and its chemical interpretation: *Geoscience Canada* 12(4), 139-146.

36. Machel H. G., Mason R.A., Mariano A.N. and Mucci A. (1991). Causes and emission of luminescence in calcite and dolomite. *Luminescence Microscopy and Spectroscopy* 25, 9-25.

37. Marshall C.R. (2006). Explaining the Cambrian "Explosion" of animals. *Annual Review of Earth and Planetary Sciences* 34, 355-384.

38. McCormick T. and Fortey R. A. (1998). Independent testing of a paleobiological hypothesis: The optical design of two Ordovician pelagic trilobites reveals their relative paleobathymetry. *Paleobiology* 24(2), 235–253.

39. Margulis L., Walker J.C.G. and Ramblerer M. (1976). Reassessment of roles of oxygen and ultraviolet light in Precambrian evolution. *Nature* 264(5587), 620-624.

40. Meyer-Rochow B.V. (2009). *Bioluminescence in Focus: A collection of illuminating essays*. Research Signpost, Trivandrum, India.

41. Nakamura H., Shirakawa Y., Kitamura H., Sato N., Shinji O. and Takahashi S. (2013). Mechanism of wavelength conversion in polystyrene doped with benzoxanthene: emergence of a complex. *Scientific Reports* 3: 2502.

42. Pedone, V. A., Cercone, K. R. and Burruss, R. C. (1990). Activators of photoluminescence in calcite: evidence from high-resolution, laser-excited luminescence spectroscopy. *Chemical Geology* 88(1), 183-190.

43. Qiu J. (2014). Oxygen fluctuations stalled life on Earth. *Nature News* (11 Juli 2014).

44. Schoenemann B. and Clarkson E.N.K. (2011). A light guide lens in ancient trilobites? *Earth and Environmental Science Transactions of the Royal Society of Edinburgh* 102(1), 17-23.

45. Schoenemann B. and Clarkson E.N.K. (2013). Discovery of some 400 million year-old sensory structures in the compound eyes of trilobites. *Scientific Reports* 3, 1429.

46. Schoenemann B., Clarkson E.N.K., Ahlberg P. and Dies Alvarez M. E. (2010). A Tiny Eye Indicating a Planktonic Trilobite. *Palaeontology* 53(4), 695-701.

- 610 47. Schindler T., Sutcliffe O., Bartels C., Poschmann M. and Wuttke M. (2002).
611 Lithostratigraphical subdivision and chronostratigraphical position of the middle Kaub
612 Formation (Lower Emsian, Lower Devonian) of the Bundenbach area (Hunsrück, SW Germany).
613 *Metalla* (Bochum) 9(2), 73-104.
614
615 48. Shimomura O. (1992): The role of superoxide dismutase in regulating the light emission of
616 luminescent fungi. *Journal of Experimental Botany* 43(11), 1519–1525.
617
618 49. Shimomura O. (2005): The discovery of aequorin and green fluorescent protein. *Journal of*
619 *Microscopy* 217, 3–15.
620
621 50. Sparks J.S., Schelly R. C.; Smith W.L., Davis M.P., Tchernov D., Pieribone V.A. and Gruber D.
622 F. (2014). The Covert World of Fish Bioluminescence: A Phylogenetically Widespread and
623 Phenotypically Variable Phenomenon. *PLoS ONE* 9 (1): e83259.
624
625 51. Struve W. (1990) Paläozoologie III (1986-1990). In *Wissenschaftlicher Jahresbericht*
626 *1988/1989 des Forschungsinstituts Senckenberg, Frankfurt am Main, Courier Forschungsinstitut*
627 *Senckenberg* 127(1990), 251–279.

628

- 629 52. Tanaka G., Schoenemann B., El Hariri K., Ono T., Clarkson E.N.K. and Maeda H. (2015).
630 Vision in a Middle Ordovician trilobite eye. *Palaeogeography, Palaeoclimatology, Palaeoecology*
631 433, 129-139.

632
- 633 53. Tortello M. F. and Esteban S. B. (2003). Lower Ordovician stratigraphy and trilobite faunas
634 from the southern Famatina Range, La Rioja, Argentina. *Special Papers in Palaeontology* 70,
635 213-239.

636
- 637 54. Towe K.M. (1973). Trilobite eyes: calcified Lenses *in vivo*. *Science* 179(4077), 1007-1009.

638
- 639 55. Wilmot N. V. (1990). Biomechanics of trilobite exoskeletons. *Palaeontology* 33(4), 749-768.
640
- 641 56. Wilmot N. V. & Fallick A. E. (1989). Original mineralogy of trilobite exoskeletons.
642 *Palaeontology* 32(2), 297-304.

643
- 644 57. Wozniak B. and Dera J. (2007). *Light Absorption in Sea Water*. Springer Science+Business
645 Media, LLC, New York.

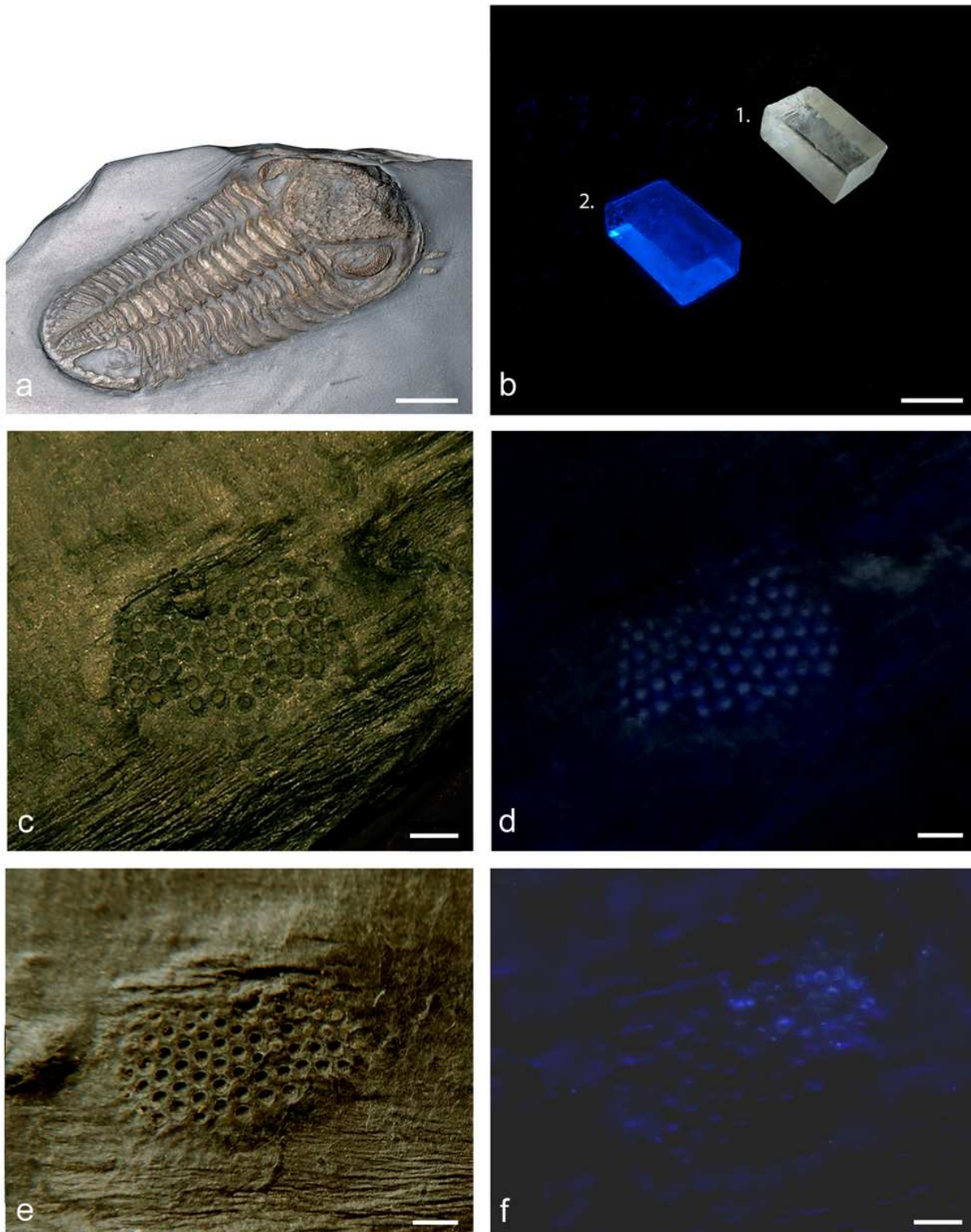
646

58. Xia J., Ito E. and Engstrom D.R. (1997). Geochemistry of ostracode calcite: Part 1. An experimental determination of oxygen isotope fractionation. *Geochimica et Cosmochimica Acta* 61(2), 377-382.

1

The glow in the calcitic lenses of a phacopid trilobite's eye.

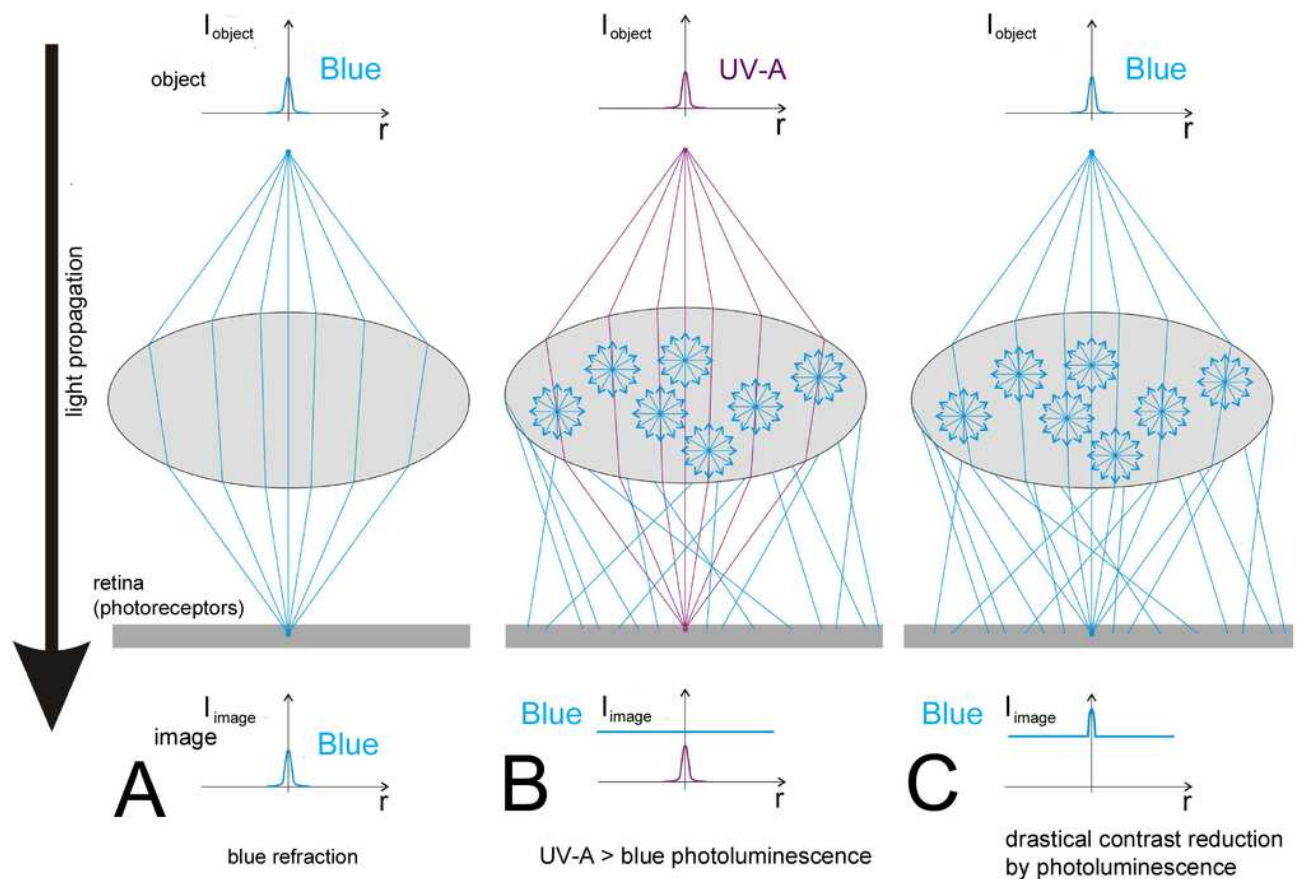
(a) *Chotecops ferdinandi* (Kayser, 1880), Bundenbachschiefer, Lower Devonian, Location: Grube Eschenbach, Hunsrück, Germany, scale bar ~1cm. (housed in the collection of Steinmann Institute, University of Bonn [still open, curator on field work]) (b) 1. Calcite crystal (~3cm), 2. Fluorescent when illuminated with ~365nm under water. (c) Isolated moult of a *Chotecops* compound eye with lenses preserved [GIK 2118]. (d) The same showing fluorescence in the calcitic lenses of the trilobite compound eye when illuminated with UVA-light (~365nm). (e) Isolated moult of a *Chotecops* compound eye with lenses preserved [GIK 2119]. (f) The same showing fluorescence in the calcitic lenses of the trilobite compound eye when illuminated with UVA-light (365nm). b-f) scale bar ~1mm.



2

The optical problem caused by the UV-A-induced photoluminescent diffuse blue light in the image formation by a dioptric apparatus.

For explanation see text



3

Underwater downward irradiance and fluorescence of trilobite lenses under UV-A light and day-light conditions.

Why the UV-A-induced photoluminescent blue-green glow in trilobite eyes and exoskeletons did not cause problems for trilobites? (a) Underwater downward irradiance (changed and simplified after Wozniak & Dera 2007) (b) Isolated moult of a *Chotecops* compound eye with lenses preserved [GIK 2118] showing a very slight fluorescence in the calcitic lenses of the trilobite compound eye when illuminated with UVA-light (~365nm) under day-light conditions. (c) Isolated moult of a *Chotecops* compound eye with lenses preserved [GIK 2119] showing a very slight fluorescence in the calcitic lenses of the trilobite compound eye when illuminated with UVA-light (365nm) under day-light conditions, but not as evident as in b). b, c) scale bar ~1mm.

