At what stratigraphical level is the mid Ludfordian (Ludlow, Silurian) positive carbon isotope excursion in the type Ludlow area, Shropshire, England?

David K. Loydell & Jiří Frýda

The balance of evidence suggests that the mid Ludfordian positive carbon isotope excursion (CIE) commences in the Ludlow area, England in the uppermost Upper Whitcliffe Formation, with the excursion continuing into at least the Platschisma Shale Member of the overlying Downton Castle Sandstone Formation. The Ludlow Bone Bed Member, at the base of the Downton Castle Sandstone Formation has previously been considered to be of Přídolí age. Conodont and thelodont evidence, however, are consistent with the mid Ludfordian age proposed here. New δ¹³C values are presented from Weir Quarry, W of Ludlow, showing a pronounced positive excursion commencing in the uppermost Upper Whitcliffe Formation, in strata with a palynologically very strong marine influence. Elsewhere in the world, the mid Ludfordian positive CIE is associated with major facies changes indicated shallowing; the lithofacies evidence from the Ludlow area is consistent with this. There appears not to be a major stratigraphical break at the base of the Ludlow Bone Bed Member. • Key words: Silurian, Ludlow, carbon isotopes, conodonts, chitinozoans, thelodonts, stratigraphy.


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It is now generally recognised that the Silurian was a period of profound global environmental changes which are reflected in the geological record by extinction events, major facies changes and positive carbon isotope excursions. In this paper an attempt is made to constrain stratigraphically the mid Ludfordian (Ludlow) positive carbon isotope excursion (CIE) in Shropshire, England, on the margins of the early Palaeozoic Welsh Basin.

This mid Ludfordian positive CIE is the largest in the Phanerozoic, with δ¹³C values exceeding +11‰, recorded from Sweden, on the palaeocontinent Baltica (Wigforss-Lange 1999), and +12–13‰ from Australia (Andrew et al. 1994), part of the Gondwana supercontinent. Given that earlier Silurian positive carbon isotope excursions have been identified in the Welsh Basin and/or on its margins (e.g. the early Sheinwoodian excursion: Loydell & Frýda 2007, Cramer et al. 2010; the mid Homerian excursion: Corfield et al. 1992), it would seem reasonable to expect that evidence for the mid Ludfordian event should be preserved in strata in the region also. Thus far, however, it has not been recognised; indeed there are no published (or, to the authors’ knowledge, unpublished) studies of the isotopic record in the Ludfordian strata of Wales and the Welsh Borderland.

The paper commences with discussions of Ludfordian biostratigraphy, both globally and in the Anglo-Welsh area in an attempt to constrain the bio- and lithostratigraphical level at which the isotope excursion should be sought. Facies changes associated with the excursion in Baltica, peri-Gondwanan Europe and Laurentia are discussed with a view to determining, from a sequence stratigraphical viewpoint, what might be expected from Shropshire’s geological record at the excursion level. Having identified the two stratigraphical levels which fulfil some or all of the necessary criteria to have the potential to preserve the excursion, the results of δ¹³C analyses (for methodology see Loydell & Frýda 2007) on samples from two carefully chosen sections are presented and discussed.

The term mid Ludfordian is preferred here to late Ludfordian (as used previously by several authors, e.g. Munnecke et al. 2003, Jeppsson 2005, Stricanne et al. 2006, Loydell 2007, Eriksson & Calner 2008) as it more accurately reflects the stratigraphical level of the excursion, as discussed below. Kaljo & Martma (2006), Barrick

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East European Platform, Poland  |  Barrandian, Czech Republic
---|---
Uncinatograptus spinifer  |  Bohemograptus praecornutus
Uncinatograptus protospinifer  |  Pseudomonoclimacis latilobus/
Uncinatograptus acer  |  Bohemograptus tenuis
Pseudomonoclimacis latilobus/
Slovinograptus balticus  |  Bohemograptus praecornutus
Neocucullograptus kozlowski  |  Bohemograptus tenuis
Neocucullograptus inexpectatus  |  Bohemograptus praecornutus
Neolobograptus auriculatus  |  Cucullo. aversus/
Bohemograptus cornutus  |  S. leintwardinensis
Slovinograptus balticus  |  Saetograptus lineolaris
Pseudomonoclimacis latilobus/
Slovinograptus balticus  |  Neocucullograptus inexpectatus
Neocucullograptus kozlowski  |  Neocucullograptus kozlowski
Neocucullograptus inexpectatus  |  Neocucullograptus kozlowski
Bohemograptus leintwardinensis  |  Saetograptus lineolaris

Figure 1. Correlation of the Ludfordian graptolite biozonations of Poland and Bohemia (from Loydell in press).

et al. (2010), Slavík et al. (2010) and Manda et al. (in press) also refer to this excursion (and/or the penecontemporaneous Lau and Kozlowski events) as mid Ludfordian.

Ludfordian biostratigraphy and the mid Ludfordian positive carbon isotope excursion

Three fossil groups are widely used in Silurian biostratigraphy: graptolites, chitinozoans and conodonts. Two other groups, ostracods and thelodonts are important in many regions; their significance in the Ludlow area is considered later in this paper when the age of the Ludlow Bone Bed Member is discussed.

Ludfordian graptolite biostratigraphy and the mid Ludfordian positive carbon isotope excursion

A remarkably detailed graptolite biozonation has been erected for the Ludfordian of the East European Platform of Poland (Baltica) with ten biozones recognised by Urbanek & Teller (1997). In Bohemia (peri-Gondwanan Europe), fewer (six) biozones are recognised. Correlations are indicated in Fig. 1. Two major extinction events have been recognised in the Ludfordian and these have a profound impact on the nature of graptolite assemblages (Urbanek 1993, Melchin et al. 1998). By the end of the earliest Ludfordian Saetograptus leintwardinensis Zone, retiolitids, Saetograptus and Cucullograptus, all important constituents of Gotarian graptolite faunas, were extinct. Urbanek (1993) referred to this major extinction event as C2 (crisis 2) or the leintwardinensis Event. Post-leintwardinensis Biozone graptolite assemblages are dominated by Neocucullograptus and Bohemograptus until both of these genera became extinct at the end of the Neocucullograptus kozlowski Zone in the second of the Ludfordian graptolite crises, C3, or the kozlowski Event.

In Kosov Quarry Bohemia, the mid Ludfordian excursion commences just above the LADs of Bohemograptus and Neocucullograptus kozlowski (Lehnert et al. 2007) and thus the excursion coincides with the kozlowski Event. This parallels the relationship between excursions and graptolite extinctions seen earlier in the Silurian (Loydell 2007).

Graptolites in the Ludlow area

Despite the absence of graptolites above the Upper Leintwardine Formation, they still provide an extremely useful biostratigraphical constraint upon the level at which to search for the mid Ludfordian excursion. The presence of the lower Ludfordian biozonal index graptolite Saetograptus leintwardinensis in the Lower and Upper Leintwardine formations (Holland et al. 1963) indicates clearly that the mid Ludfordian excursion, if recorded in the rock record, must lie at a higher stratigraphical level.

Ludfordian chitinozoan biostratigraphy and the mid Ludfordian positive carbon isotope excursion

Verniers et al. (1995), in their ‘global Chitinozoa biozonation for the Silurian’, recognised two Ludfordian biozones. The lower, Eisenackitina philipi Biozone has its base defined by the FAD of the eponymous species in the Ludlow area. The succeeding chitinozoan biozone is the E. barrandeii Biozone. An unzoned interval was placed between these two biozones, reflecting the paucity of chitinozoans above the LAD of E. philipi in the Ludlow area and general lack of studies at this time elsewhere on this interval.

There is a consistent relationship between the FADs of Eisenackitina philipi and E. barrandeii: in all published sections the FAD of the former occurs several tens of metres below that of the latter. Although used successfully as a biozonal index in southern Sweden (Grahn 1996), in the East Baltic region E. philipi is rare, prompting Nestor (2009) to erect an E. lagenomorpha Biozone for the lower Ludfordian. The FAD of confidently identified E. philipi in the East Baltic region is at the base of, or very low in the E. lagenomorpha Biozone (Nestor 2009), suggesting that the bases of the E. philipi and E. lagenomorpha biozones are at an approximately synchronous level. It appears that the LAD of E. philipi is at a lower stratigraphical level than that of E. lagenomorpha: this is certainly the case on Gotland (Laufeld 1974) and in the East Baltic (Nestor 2009).

The stratigraphical utility of other late Silurian chitinozoan taxa is highly variable. Urnochitina gr. urna is
stated by Paris (in Kříž et al. 1986) to be ‘most probably completely restricted to the Přídolí’ and this is obviously therefore an important and useful species group, particularly as several studies have recorded chitinozoans through the uppermost Ludlow/lower Přídolí transition. Sphaerochitina sphaerocephala has been consistently recorded only from the E. barrandei Biozone and higher strata and also therefore appears to be an important species, despite Laufeld’s (1974) misgivings that “it is hard to escape the conclusion that S. sphaerocephala has become a waste-basket taxon”. The level of first appearance of S. sphaerocephala is, however, difficult to assess: in several sections it is present in the lowest sample studied, e.g. the Požáry and Koledník quarries, Bohemia (Kříž et al. 1986), the A1-61 borehole, Libya (Jaglin & Paris 2002) and thus may well occur in lower strata. On Gotland, from which E. barrandei has not been recorded, the FAD of S. cf. sphaerocephala at the base of the Hamra Formation is immediately above the Burgsvik Formation, a unit with an unusual chitinozoan assemblage, considered to be highly facies dependent (Laufeld 1974) and thus the absence of S. cf. sphaerocephala may be environmentally rather than stratigraphically controlled. The ‘cf.’ used by Laufeld (1974) reflects the ‘atypical’ appearance of the lowest Hamra Formation specimens (from unit a, which is 0.2–1.5 m thick out of a total formational thickness of 40 m). S. sphaerocephala is a highly variable species (see e.g. Wrona 1980, fig. 9), so it may be that these atypical specimens can be accommodated within the species. Typical S. sphaerocephala were found by Laufeld (1974) in unit b of the Hamra Formation. Jeppsson et al. (2006) assigned the Hamra Formation to the Ozarkodina snajdri Biozone. The lower part of this formation, including unit b, is still within the mid Ludfordian excursion (Stricanne et al. 2006). In the Klintabomminger 1 section, Sweden, the FAD of S. sphaerocephala is immediately above the LAD of E. philipi within the long stratigraphical ranges of E. lagenomorpha and Angochitina echinata; E. barrandei was not identified from this section (Grahn 1996). The available evidence, particularly that from Gotland, suggests that the stratigraphically earliest S. sphaerocephala appeared significantly before the first E. barrandei. This is important with regard to the dating of the mid Ludfordian excursion in the Ventspils D-3 core, Latvia (see below).

As knowledge of chitinozoan biostratigraphy has increased, other taxa, previously thought to be of stratigraphical importance, have been shown to have longer ranges than originally thought and these cannot therefore be used with any confidence in high resolution studies. For example, Pterochitina perivelata is part of Paris’s (in Kříž et al. 1986) Přídolí assemblage, but has recently been found by Nestor (2009) throughout much of the Ludfordian of the Pavilosta core, Latvia. Similarly, Anycrochitina pedavis, stated to be ‘restricted to the uppermost Ludlow’ by Paris (in Kříž et al. 1986), occurs in both the E. lagenomorpha and E. barrandei biozones of the Pavilosta core (Nestor 2009).

**Chitinozoan dating of the mid Ludfordian excursion**

In the Ventspils D-3 core the mid Ludfordian excursion occurs between depths of 470 m and 460 m (Kaljo et al. 1998). According to Nestor (2009, fig. 4) this level is within the lower part of the Eisenackitina barrandei Biozone. The biozonal species, however, was not confidently identified in the core. Dating of the CIE in the Ventspils D-3 core as E. barrandei Biozone hinges on Sphaerochitina sphaerocephala, confidently identified immediately below the excursion from the –472 m sample, not occurring below the E. barrandei Biozone. The stratigraphical significance of this species is discussed at length above and it is concluded, however, that its first appearance was most probably before that of E. barrandei. The FAD of confidently identified E. philipi in the Ventspils D-3 core is approximately 40 m below the onset of the excursion; its LAD is within the excursion interval. The isotope excursion occurs at a level (upper Mituva Formation) at which chitinozoan diversity declined markedly and in the overlying Ventspils Formation chitinozoans are ‘scarce’. Nestor (2009) related the diversity decline to the Lau Event and refers to ‘intense extinction of chitinozoans’. A major extinction at this stratigraphical level is not apparent from the data from the other cores (Ohesaare, Pavilosta and Dubovskoye) studied by Nestor, however, nor from her summary range chart (fig. 7) suggesting that local environmental/preservational factors are responsible for the reduced chitinozoan diversity and abundance seen in the Ventspils D-3 core.

A peak $\delta^{13}$C$_{carb}$ value of 4.2‰ was recorded in the Pavilosta core, Latvia by Kaljo et al. (1998). This was at a depth of 739 m (Kaljo & Martma pers. comm., December 2010), within the lower half of the Eisenackitina lagenomorpha Biozone and 44 m below the FAD of Eisenackitina barrandei.

In both cores (Ventspils D-3 and Pavilosta) the excursion occurs significantly above the FAD of Eisenackitina philipi. The LAD of E. philipi lies 3 m below the peak $\delta^{13}$C$_{carb}$ value in Pavilosta and within the excursion interval in the Ventspils D-3 core. On Gotland the LAD of E. philipi also lies within the excursion, in the Burgsvik Formation (Laufeld 1974).

Isotope and chitinozoan data are available also from the Czech Republic. In the section close to the tunnel entrance at Požáry Quarry the mid Ludfordian CIE has been recorded in beds 34–41 (Lehnert et al. 2007), Eisenackitina barrandei was recorded by Paris & Kříž (1984) from beds 87–96 (highest Ludlow to lowermost Přídolí; note that beds below bed 87 were not studied). Bed 87 is more than
10 m above the excursion. This indicates that the excursion is not within the upper part of the *E. barrandei* Biozone and indeed need not lie within this biozone at all.

### Chitinozoans in the Ludlow area

A thorough review of the Ludlow chitinozoans of the Ludlow area was provided by Sutherland (1994). As noted above, the base of the *E. philipi* Biozone was defined in the Ludlow area (Verniers *et al.* 1995). The biozone here encompasses the upper part of the Upper Leintwardine Formation, the entire Lower Whitcliffe Formation and lower part of the Upper Whitcliffe Formation. Above this level chitinozoans are rare, but still present in the upper part of the Upper Whitcliffe Formation (Richardson & Rasul 1990). Sutherland (1994) explained this by suggesting that during the Ludfordian increased input of fresh water dramatically reduced the numbers of chitinozan producers living in the shelf seas surrounding the Welsh Basin.

Richardson & Rasul (1990, fig. 10b) illustrated a chitinozan, identified as *E. barrandei* Paris & Kříž, 1984 by Florentin Paris, from their sample 161 in the Upper Whitcliffe Formation. This sample is from 0.23–0.26 m below the base of the Ludlow Bone Bed Member of the Downton Castle Sandstone Formation at Weir Quarry, near Downton ca 5 km W of Ludlow. The identification of *E. barrandei* was accepted by Verniers *et al.* (1995) in their global biozonation paper. Miller *et al.* (1997) subsequently erected new chitinozan species from Nantyrhynau Quarry, eastern Wales, more than 30 km WNW of Weir Quarry, in deeper water, turbiditic strata, considered to straddle the Ludlow/Přídlí series boundary and be the lateral equivalent of the upper part of the Upper Whitcliffe Formation and lower Downton Castle Sandstone Formation. One of their new species, *Eisenackitina clunensis*, is very similar to *E. barrandei*, differing only in the position of vesicle flexure. In *E. clunensis* this occurs at a mid-point on the vesicle flanks, whereas in *E. barrandei* the flexure is closer to the aperture (Miller *et al.* 1997). This difference is significant now that Nestor has recorded all of Miller *et al.*’s (1997) new species from the East Baltic region and shown that all of them here have different or longer stratigraphical ranges than seen in Nantyrhynau Quarry. Of particular significance is the occurrence of *E. clunensis* in the upper Angochitina elongata and lower *Eisenackitina lagenomorpha* biozones, with a LAD below that of *E. philipi*. In combination with the Welsh data, this indicates that this species has a long stratigraphical range, and probably it cannot be used in high resolution biostratigraphy. Why is this significant? In Richardson & Rasul’s (1990) specimen from the Upper Whitcliffe Formation of Weir Quarry the location of flexure suggests that this specimen is not in fact *E. barrandei*, but instead is *E. clunensis*. Thus the Upper Whitcliffe Formation/Downton Castle Sandstone Formation boundary need not be at a stratigraphical level above the base of the *E. barrandei* Biozone (i.e. uppermost Ludfordian).

### Ludfordian conodont biostratigraphy and the mid Ludfordian positive carbon isotope excursion

Although Corradini (2009) emphasized that much work remains to be done before there is a widely accepted upper Silurian ‘Standard Conodont Zonation’, all six of the conodont biozonations for the Ludfordian presented were united in having the lower Ludfordian Ancoradella ploeckensis Biozone succeeded by a Polygnathoides siluricus Biozone. Similarly, all six terminated the Ludfordian Stage with the *Ozarkodina crispa* Biozone. The greatest variation lay between the *siluricus* and *crispa* biozones with *P. latialata* and/or *O. snajdri* biozones recognised. Jepsson (2005) erected an Icriodontid Biozone for this interval. Conodont occurrence data provided by various authors (e.g. Miller 1995, Koren’ & Walliser 1998, Viira & Aldridge 1998, Corradini & Serpagli 1999) indicate that conodont dating and correlation of Ludfordian strata may not, however, be as straightforward as published charts of conodont biozones might imply.

In this respect, it is interesting to compare the chitinozan data of Nestor (2009) for the Ohesaare core, Saaremaa (Estonia) with the conodont biozonation of Viira & Aldridge (1998, fig. 6). Here the *O. crispa* Biozone is recognised from −95.95 m to −101.05 m. According to all of the biozonations shown by Corradini (2009), the *O. crispa* Biozone is the highest in the Ludfordian, yet the chitinozoans within the stratigraphical range of *O. crispa* in the Ohesaare core indicate the uppermost Angochitina elongata and lower *Eisenackitina lagenomorpha* biozones, not the upper Ludfordian *Eisenackitina barrandei* Biozone. The highest Ludfordian conodont biozone in the Ohesaare core is the *Ozarkodina parasnajdri* Biozone. Lithostratigraphical and thelodont evidence also point to a stratigraphical level for the Saaremaa *O. crispa* substantially lower than the upper Ludfordian (possibly even upper Gorstian). Viira & Aldridge (1998, p. 48) concluded that: “An explanation may be that the fluctuating nearshore environment of Saaremaa was suited to the earlier evolution of populations of *O. crispa.*”

Viira & Aldridge (1998) distinguished three morphs of *Ozarkodina crispa*. The stratigraphically early specimens from Ohesaare (and several other cores and outcrops) belong to morphs α₁ and α₃. The stratigraphical significance of morph α₃ in the Ludlow area is discussed below.

Jepsson’s (2005) revision of the Ludfordian conodont biostratigraphy for Gotland enables the isotope excursion...
here to be dated precisely: the excursion commences in the uppermost part of the *P. siluricus* Biozone; high values continue through the Icriodontid Biozone and much of the *O. snajdri* Biozone; they decline towards the top of the *O. snajdri* Biozone (Stricanne *et al.* 2006, fig. 3). Kaljo & Martma (2006) showed the relationship of the mid Ludfordian excursion to biostratigraphical data from several fossil groups. For the Vidukle and Ventspils cores, the pattern is consistent with that on Gotland, with the LAD of *P. siluricus* just below the excursion. In Ventspils, *O. snajdri* occurs within the excursion interval (Kaljo & Martma 2006, fig. 4), but as this species has a long stratigraphical range (appearing in the lower Gorstian; Jeppsson 2005, p. 276), the stratigraphical significance of this occurrence is limited.

In Bohemia (peri-Gondwanan Europe) the beginning of the excursion has also been dated to the *P. siluricus* Biozone (Lehnert *et al.* 2007, Slavík *et al.* 2010) as it has also in the United States (Barrick *et al.* 2010). In the latter no Icriodontid Biozone was recognizable.

**Conodonts in the Ludlow area**

Miller (1995) recorded conodonts identified as *O. cf. crispa* from the Upper Whitcliffe Formation 0.3–0.15 m below the Ludlow Bone Bed Member at Ludford Corner, Ludlow. Viira & Aldridge (1998, p. 48) identified these as their morph α3, Miller (1995, p. 374) used the presence of *O. cf. crispa* in the uppermost Upper Whitcliffe Formation at Ludford Corner as evidence that the “base of the Přídolí Series in Britain is very close to the level of the base of the Downton Castle Sandstone Formation”. The identification of the material by Viira & Aldridge (1998) as their morph α3, and the stratigraphical level at which this morph occurs on Estonia suggest that the uppermost Upper Whitcliffe Formation in Ludlow could be considerably older than latest Ludfordian. The significance of this is discussed below.

Jeppsson & Aldridge (2000), whilst noting that in the Ludlow area “conodont information is sparse”, suggested that the Lau Event (a conodont extinction event stratigraphically within the mid Ludfordian positive CIE on Gotland) “is probably represented close to the boundary between the Upper Leintwardine Formation and the Lower Whitcliffe Formation”.

**Facies changes and unconformities associated with the mid Ludfordian positive carbon isotope excursion**

The mid Ludfordian positive CIE is associated with major facies changes indicating shallowing and/or with a stratigraphical break. This shallowing is reflected in Johnson’s (1996) eustatic sea-level curve for the Silurian where an interval of low sea-level is shown in the mid Ludfordian (between highstands 7 and 8).

Not surprisingly, facies changes indicating shallowing have been recorded from many sections worldwide. On western Gotland (Baltica) the excursion commences close to the boundary between the Hemse Group (distal shelf mudstones with graptolites) and the Eke Formation (which includes proximal shelf oncolitic limestones) (Jeppsson 2005, Stricanne *et al.* 2006; see Eriksson & Calner 2008 for a more detailed facies analysis). At this stratigraphical level on eastern Gotland, Cherno’s (1982) recognised palaeokarsts indicating emergence; Jeppsson (2005), Jeppsson *et al.* (2007) and Eriksson & Calner (2008) also recognised a ‘brief hiatus’ at this level. A very similar shallowing trend to that on Gotland, from marls to oncolidal limestones, was recorded from Australia (Gondwana) by Jeppsson *et al.* (2007).

In Kosov Quarry, Bohemia (peri-Gondwanan Europe), graptolitic strata below the CIE indicating a ‘deeper and less oxygenated’ environment are overlain by shallower marine, thicker bedded limestones, in which the excursion is recorded (Lehnert *et al.* 2007); in Požáry Quarry, Bohemia the excursion occurs in strata exhibiting ‘rapid shallowing’ with massive limestone beds with palaeokarst development (Lehnert *et al.* 2007); a gap in the conodont record has recently been recognised at this level (Slavík *et al.* 2010). Palaeokarst development at the same
greater duration and thus no major positive isotope excursion. A graphical break within the Ludfordian appears to be of lesser duration in the Ohesaare core, Estonia (Baltica), however, the stratigraphical sections both record the CIE, albeit incompletely. In the biostratigraphical viewpoint, the level of the stratigraphical transition is discussed first.

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Where is the mid Ludfordian carbon isotope excursion in the Ludlow area?

Based on the above stratigraphical and facies observations, one would expect the mid Ludfordian positive CIE in the Ludlow area succession to be present above the LAD of S. leintwardinensis (in the Upper Leintwardine Formation) and be marked by a major facies change and/or a stratigraphical break. Within the Ludlow area such a facies change occurs at one level only, between the Upper Whitcliffe Formation and Downton Castle Sandstone Formation, the base of which is marked by the Ludlow Bone Bed Member. It has been suggested previously, however, that the CIE may lie close to the Upper Leintwardine Formation/Lower Whitcliffe Formation boundary, so this stratigraphical transition is discussed first.

The Upper Leintwardine Formation/Lower Whitcliffe Formation boundary

As noted above, Jeppson & Aldridge (2000) suggested that the Lau Event (a conodont extinction event stratigraphically close to the mid Ludfordian positive CIE) “is probably represented close to the boundary between the Upper Leintwardine Formation and the Lower Whitcliffe Formation”.

To see whether the excursion could be recognised at this level, samples were collected through the uppermost 2 m of the Upper Leintwardine Formation (nodular calcareous siltstones) and lowermost 4 m of the Lower Whitcliffe Formation (calcareous siltstones) from newly quarried sections at Delbury, Shropshire (SJ 494 862; see Fig. 2 for location). δ¹³C<sub>carb</sub> analyses indicate that no major positive CIE is present within this stratigraphical interval (Fig. 3). The oxygen isotope data (Table 1) indicate a diagenetic influence on the results, but, despite this, the carbon results should display some evidence for a CIE of the magnitude of the mid Ludfordian excursion were it to be present. Isotope data from many other Silurian sections exhibit very low oxygen values (below –10‰), indicative of diagenetic modification of the oxygen isotope record, but preserve a largely unmodified carbon isotope signal (see e.g. data.
presented in Andrew et al. 1994, Wigforss-Lange 1999, Cramer et al. 2006) with excursions clearly recognizable. One might argue that it is possible that there is no CIE in this interval due to isolation of the Welsh Basin resulting from progressive closure of the Iapetus Ocean (as suggested by Woodcock et al. 1988, Smith & Ainsworth 1989). However, this is unlikely because these strata (and those overlying them) contain geographically widespread conodont, thelodont and chitinozoan taxa (e.g. Miller 1995, Miller et al. 1997, Miller & Märs 1999), suggesting good connections with the open ocean.

Chitinozoan biostratigraphy indicates that this is not the level to look for the mid Ludfordian positive CIE anyway. The Upper Leintwardine Formation/Lower Whitcliffe Formation boundary is low within the Eisenackitina philipi Biozone. As noted above, where dated with chitinozoans the excursion is just above or includes the LAD of *E. philipi*.

### The Upper Whitcliffe Formation/Downton Castle Sandstone Formation boundary

The base of the Downton Castle Sandstone Formation is marked by the Ludlow Bone Bed Member. This is succeeded by the Platyschisma Shale Member which in turn is overlain by the Sandstone Member.

Richardson & Rasul (1990) conducted a detailed palynofacies analysis through the Upper Whitcliffe Formation and lower Downton Castle Sandstone Formation at Weir Quarry, Downton (SO 45607525). For this reason, this locality was chosen for sampling for isotopic analysis as comparison of isotope results would be possible with both lithological and palynological changes. The very low carbonate content and dolomitization precluded meaningful analysis of δ¹³C_carb from the samples; the presence of well preserved organic matter, however, enabled analysis of δ¹³C_org, the resulting curve being presented in Fig. 4.

The δ¹³C_org curve clearly shows the commencement of a major positive excursion in the uppermost part of the Upper Whitcliffe Formation, with the highest δ¹³C_org value of –23.88‰ occurring 0.78 m above the base of the Ludlow Bone Bed Member, within the Platyschisma Shale Member.

Comparison with Richardson & Rasul’s (1990) results is very interesting. The excursion commences in strata which, from a palynological perspective, are fully marine. Richardson & Rasul (1990) introduced a marine influence index (in which 0 = fully terrestrial and 100 = fully marine) based upon the composition of palynological assemblages. Samples from 19–25.5 cm and 0–5 cm below the Ludlow Bone Bed have a marine influence index of 96.5 and 97.5 respectively. A major change in palynofacies occurs at the base of the Ludlow Bone Bed Member exemplified by the increase in percentage of spores within the assemblages, from 2.5% in the highest Upper Whitcliffe Formation sample to 50.5–81.5% in the 20–30 mm thick Ludlow Bone Bed itself. The δ¹³C_org values show a rapid increase through the Ludlow Bone Bed Member and lowermost Platyschisma Shale Member, reaching –24.13‰ 5 cm above the base of the latter. The major part of the rising limb of this δ¹³C_org excursion is therefore within strata with a relatively consistent palynological signal, with spores dominant (values are between 50.5% and 81.5% throughout; Richardson & Rasul 1990, table 2) with the accompanying acritarchs predominantly sphaeromorphs and *Micrhystridium*.

It is rather surprising that the major change in palynofacies is not the level at which the δ¹³C_org values show their greatest change, given that that the organic-walled palynomorphs might be expected to give a good indication

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**Table 2.** δ¹³C_org data for Weir Quarry. Measurements (in metres) for sampling levels are taken from the base of the Ludlow Bone Bed Member.

<table>
<thead>
<tr>
<th>Sample</th>
<th>δ¹³C_org</th>
<th>Sample</th>
<th>δ¹³C_org</th>
</tr>
</thead>
<tbody>
<tr>
<td>+1.28</td>
<td>–24.18</td>
<td>–0.25 –0.26</td>
<td>–26.82</td>
</tr>
<tr>
<td>+1.03</td>
<td>–24.39</td>
<td>–0.60</td>
<td>–27.09</td>
</tr>
<tr>
<td>+0.78</td>
<td>–23.88</td>
<td>–0.75</td>
<td>–27.04</td>
</tr>
<tr>
<td>+0.53</td>
<td>–24.96</td>
<td>–1.00</td>
<td>–27.15</td>
</tr>
<tr>
<td>+0.28</td>
<td>–24.13</td>
<td>–1.10</td>
<td>–26.61</td>
</tr>
<tr>
<td>+0.15–0.19 (repeat)</td>
<td>–25.08</td>
<td>–1.50</td>
<td>–27.35</td>
</tr>
<tr>
<td>+0.15–0.19 (repeat)</td>
<td>–25.14</td>
<td>–1.50 (repeat)</td>
<td>–27.54</td>
</tr>
<tr>
<td>+0.05–0.07 (repeat)</td>
<td>–25.72</td>
<td>–1.75</td>
<td>–26.28</td>
</tr>
<tr>
<td>+0.05–0.07 (repeat)</td>
<td>–25.66</td>
<td>–2.25</td>
<td>–26.31</td>
</tr>
<tr>
<td>–0.03</td>
<td>–26.12</td>
<td>–2.25 (repeat)</td>
<td>–26.48</td>
</tr>
<tr>
<td>–0.12–0.15</td>
<td>–26.57</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
of the organic content of the sample as a whole. It is also important to note that in the lower Downton Castle Sandstone Formation the excursion is developed in marine strata, albeit with a significant terrestrial influx of palynomorphs. Richardson & Rasul’s (1990) marine influence index for the lower part of the Platschisma Shale Member (for which the $\delta^{13}$Corg data are plotted in Fig. 4) is 34.2 to 42.5 (in unworked samples).

Stricanne et al. (2006) recorded in detail the changes in palynomorph assemblages through the Ludfordian excursion interval on western Gotland. Perhaps not surprisingly, given that in both the Ludlow area and on Gotland (as elsewhere) the excursion is developed during a major regression, the Gotland palynological signal is similar to that at Weir Quarry: spores increase in both absolute and relative abundance.

In other Silurian sections studied for both $\delta^{13}$Corg and $\delta^{13}$Ccarb, whilst these may provide differing signals for individual samples, major excursions are clearly identified in both sets of data: e.g. the sedgwickii Zone (late Aeronian, Llandovery) excursion in Arctic Canada (Melchin & Holmden 2006); early Sheinwoodian excursion in Wales (Loydell & Frýda 2007, Cramer et al. 2010) and early Sheinwoodian and mid Homerian excursions in Arctic Canada (Noble et al. 2005).

Based on biostratigraphy, facies changes and sea-level changes, the most likely level for the beginning of the mid Ludfordian excursion is therefore high in the Upper Whitcliffe Formation with it continuing through at least the lower part of the Downton Castle Sandstone Formation. Several key questions then emerge. Firstly, is the $\delta^{13}$Corg signal at Weir Quarry the local manifestation of this major mid Ludfordian excursion or is it simply reflecting a change in the source of organic material, with an increase in terrestrial input resulting from shallowing? If the latter, then why does the major positive shift in carbon isotope values not occur at the level when spore abundance increases twenty-fold?

Secondly, is there a significant stratigraphical gap between the Upper Whitcliffe Formation and the Ludlow Bone Bed Member? If yes, then all or part of the mid Ludfordian excursion may not be preserved at all in strata in the Ludlow area. If there is not a significant gap, which would be suggested by comparison with sedimentation patterns in the Czech Republic and the Baltic region, then is the Ludlow Bone Bed Member of Ludfordian age rather than a local manifestation of organic matter.

**Figure 4.** Percentage spore abundance and $\delta^{13}$Corg curves through the Upper Whitcliffe Formation and Lower Downton Castle Sandstone Formation exposed at Weir Quarry (SO 45607525). Log is modified from Richardson & Rasul (1990, figs 2, 3), using their sample numbers (16G, A6, etc.). Additional data points (for spore %) are from Richardson & Rasul (1990, tab. 2). $\delta^{13}$Corg data is that from Table 2. Where two analyses (palynological or isotope) were conducted upon the same stratigraphical interval, both points are plotted. Abbreviation: B. B. = Bone Bed.
than of Přídolí age as it is generally portrayed? Some of these questions are discussed further below.

What age is the Ludlow Bone Bed Member?

In the absence of graptolites and chitinozoans and with conodonts very rare, fragmentary and abraded, making identification difficult (Miller 1995), other fossil groups must be used for dating the Ludlow Bone Bed Member (and overlying members of the Downton Castle Sandstone Formation). Ostracods and thelodonts both occur in the Downton Castle Sandstone Formation: the dates that they provide, however, are different and, in the light of the above observations regarding the chitinozoan and conodonts from the uppermost Upper Whitcliffe Formation and the lithological signature of the mid Ludfordian excursion in other regions and short duration of any stratigraphical breaks identified, detailed discussion of the ostracod and thelodont results is warranted here.

Ostracod evidence

The ostracods from the Ludlow Bone Bed Member indicate a Přídolí age (Miller 1995). This age has been widely accepted and a basal Přídolí age for the Ludlow Bone Bed Member is shown in the vast majority of (if not all) recent publications on the Silurian System.

The key ostracod species is *Frostiella groenvalliana* which has been confidently identified only from Přídolí age strata, from many locations particularly in the Baltic Region. As Miller (1995) noted, *F. groenvalliana* has been reported from a wide range of environments and its stratigraphical distribution is ‘remarkably consistent’ throughout its geographical range. Miller (1995, p. 369) concluded: “Because of its apparently wide facies tolerance, the sudden appearance of *F. groenvalliana* at the base of the Ludlow Bone Bed Member is therefore unlikely to be influenced entirely by a marked facies change at that level.”

Thelodont evidence

Märrs & Miller (2004) provide a very useful summary of thelodont occurrences in the Welsh Borderland which enables correlation with Baltic sections. The Lower Leintwardine Formation contains *Paralogania kaarmisensis* and *Phlebolepis elegans*. These taxa co-occur in the Himmite Beds of the Paadla Regional Stage in Estonia. The earliest Ludfordian graptolite *Saeograptus leintwardienensis* occurs in the Lower Leintwardine Formation, and the Himmite Beds are of a similar early Ludfordian age (Jeppsson et al. 1994, fig. 3). *Thelodus parvidens* has its FAD in the Welsh Borderland in the Whitcliffe Group; in Estonia its FAD is in the Uduvere Beds of the Paadla Regional Stage. These beds lie below the stratigraphical break between the Paadla and Kuressaare Regional Stages on Saaremaa. This is the stratigraphical break discussed above, at the level of the mid Ludfordian excursion. Finally, Märrs & Miller (2004) discuss *Paralogania ludlowiensis* which they state is restricted to the Downton Castle Sandstone Formation in the Welsh Borderland (Fig. 5). Miller & Märrs (1999) list occurrence data for this species on Gotland and in the East Baltic; in the Ohesaare core, Estonia, *Pa. ludlowiensis* occurs in the lower Kuressaare Formation, of undisputed Ludfordian age, within the *Eisenackitina lagenomorpha* Biozone (Nestor 2009); in the Ventspils D-3 core, Latvia *Pa. ludlowiensis* occurs in the upper Mituva Formation, at a level within the mid Ludfordian excursion; likewise, on Gotland it occurs in the Eke Formation within the mid Ludfordian excursion. Lithuanian occurrences of *Pa. ludlowiensis* are from higher stratigraphical levels, but are still of Ludfordian age. There are no Přídolí occurrences of *Pa. ludlowiensis* in the Baltic area.

Based on the presence of the thelodont *Paralogania ludlowiensis* not only in the Ludlow Bone Bed Member but also in the Sandstone Member of the Downton Castle Sandstone Formation and the stratigraphical occurrences of this species in Estonia and Latvia, the lower part at least of the Downton Castle Sandstone Formation is probably of mid Ludfordian age.

Reconciling the evidence

Clearly there is a problem with the biostratigraphical dating of the Ludlow Bone Bed Member. The ostracods, particularly *Frostiella groenvalliana*, indicate a Přídolí age, whilst the thelodonts (*Paralogania ludlowiensis*) indicate the Ludfordian. How can this anomaly be explained? Viira & Aldridge (1998) explained the apparently early stratigraphically records of *Ozarkodina crispa* in the Baltic region by suggesting that the species evolved here and later migrated into the Welsh Basin, where it is found in the uppermost part of the Upper Whitcliffe Formation. Presumably a similar explanation is required for the ‘early’ occurrences of *Paralogania ludlowiensis* in the Baltic region. Why could it not be that instead *Frostiella groenvalliana* evolved in the relatively restricted Welsh Basin in the Ludfordian and then spread to other regions in the Přídolí? Both scenarios are equally valid and are supported by extensive data: for example Viira & Aldridge (1998) recorded *O. crispa* in the middle Ludfordian Paadla Regional Stage from a total of 23 separate samples from five different cores and four further outcrop localities.

The ostracod dating requires the mid Ludfordian excursion and associated major facies change to occur between
the Baltic region based upon the conodont provided by correlation with the Paadla Regional Stage of morph crispa and the thelodont as the major global environmental changes associated recorded at the formational boundary is the local manifestation and the Ludfordian age and that the marked lithological change recorded in the Ludlow Bone Bed Member, is much more likely to be taken from Siveter et al. (1989, fig. 30). Note that the stratigraphical range of Saetograptus leintwardinensis is as shown in Holland et al. (1963): the species may not range from the base of the Lower Leintwardine Formation to the top of the Upper Leintwardine Formation, but does occur in both formations. Note also that the LAD of Eisenackitina philipi is at a level at which chitinozoans become extremely sparse in the Ludlow area. Abbreviation: U. L. = Upper Leintwardine.

the Saetograptus leintwardinensis-bearing beds in the Upper Leintwardine Formation and the Ludlow Bone Bed Member. The only candidate level, the Upper Leintwardine/Lower Whitcliffe Formation transition, has no associated isotope excursion and chitinozoan data suggest that this is too low a stratigraphical level from a biosociation of the East Baltic region. Aldridge (1998) in an attempt to explain the apparently anomalous conodont records which does not require a significant stratigraphical break beneath the Ludlow Bone Bed Member comes from the nature of stratigraphical breaks in other sections recording the mid Ludfordian excursion. In all cases, as discussed above, the breaks are of very limited stratigraphical extent, with even the most substantial break, at Ohesaare, Estonia, being contained within part of a single chitinozoan biozone.

A shorter stratigraphical break was proposed by Viira & Aldridge (1998) in an attempt to explain the apparently anomalous conodont records of the East Baltic region when compared with the Welsh Borderland. As indicated above there is an alternative explanation for these anomalous conodont records which does not require a significant stratigraphical break in the Ludlow area succession. Also the carbon isotope curve derived from the closely spaced samples through the Upper Whitcliffe Formation/Ludlow Bone Bed Member transition does not show any marked inflection at the formational boundary, as might be expected if there was a significant stratigraphical break at this level. We are therefore in agreement with Bassett et al. (1982) in considering that any stratigraphical break at this level represents only a very small (certainly not of the order of one biozone) amount of time.

Figure 5. Summary stratigraphy of the Ludfordian of the Ludlow area, with key stratigraphical ranges/occurrences of taxa indicated. Lithologies taken from Siveter et al. 1989, fig. 30. Note that the stratigraphical range of Saetograptus leintwardinensis is as shown in Holland et al. (1963): the species may not range from the base of the Lower Leintwardine Formation to the top of the Upper Leintwardine Formation, but does occur in both formations. Note also that the LAD of Eisenackitina philipi is at a level at which chitinozoans become extremely sparse in the Ludlow area. Abbreviation: U. L. = Upper Leintwardine.

A similar conclusion regarding the age of the Upper Whitcliffe Formation/Ludlow Bone Bed Member boundary has been reached previously (e.g. Schönlaub in Küh et al. 1986, Aldridge & Schönlaub 1989), prior to publication of much of the evidence presented above and the widespread acceptance of the ostracod dating of the Ludlow Bone Bed Member as Přídolí.

Is there a major stratigraphical break between the Upper Whitcliffe Formation and the Ludlow Bone Bed Member?

The base of the Ludlow Bone Bed Member marks an abrupt lithological change from the underlying Upper Whitcliffe Formation. Palynofacies also change markedly at this level (Richardson & Rasul 1990). Almost certainly there was an interval of non-deposition and/or erosion at the boundary between the two formations: there has been some debate as to how much time this interval represents.

Bassett et al. (1982) concluded that there was “no evidence of a measurable time break” between the Upper Whitcliffe Formation and Ludlow Bone Bed Member. This was based on the “continuity of marine deposition across the boundary”, the sedimentological evidence suggesting “merely slow or locally interrupted deposition” and the fact that the “ranges of numerous species of shelly macrofossils, thelodonts, ostracodes, conodonts, miospores and chitinozoans span the boundary” (although subsequent work has indicated otherwise for some groups). Further support for the likely short duration of any stratigraphical break beneath the Ludlow Bone Bed Member comes from the nature of stratigraphical breaks in other sections recording the mid Ludfordian excursion. In all cases, as discussed above, the breaks are of very limited stratigraphical extent, with even the most substantial break, at Ohesaare, Estonia, being contained within part of a single chitinozoan biozone.
Conclusions

With the exception of the presence of the ostracod *Frostiella groenvalliana* in the Ludlow Bone Bed Member, all data (conodont, thelodont, lithological, sequence stratigraphical and carbon isotopic) are consistent with the Ludlow Bone Bed Member being of mid Ludfordian age and the Upper Whitchcliffe Formation/Ludlow Bone Bed Member transition being a reflection of the major environmental changes associated with the mid Ludfordian positive CIE which we record here for the first time in the Ludlow area (and indeed from Britain) from Weir Quarry, with the commencement of the excursion high in the Upper Whitchcliffe Formation.

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