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Finding of trout (*Salmo* cf. *trutta*) in the Northern Jordan Valley (Israel) at the end of the Pleistocene: Preliminary results

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ABSTRACT

Tens of thousands of fish bones were recovered from the Final Natufian (Late Epipaleolithic) layer of the site of Eynan/Ain Mallaha (Northern Israel) dated to the end of the Pleistocene. Almost a hundred of them were attributed to a Salmoninae, most probably a trout *Salmo* cf. *trutta*. This is the southernmost attestation of a *Salmo* species in the Near East, past and present. It is suggested that trout were present locally as small populations at least at the end of the Pleistocene.

1. Introduction

The mid-1960s witnessed a growing awareness of the widespread importance of fish for the diet of late prehistoric societies. This trend was fuelled by influential publications such as those by Lee (1979) and Binford (1968). Lee underlined that, according to ethnographic records, fish are often the main food resource in temperate latitudes (Lee, 1979: 42) while Binford suggested a stress model where: “increased dependence on aquatic resources during the terminal and immediately post-Pleistocene period” was supposed to induce an increase in sedentism, population growth, food storage and finally grain agriculture (Binford, 1968: 332–333). Thus, an intellectual stimulus was set down to prompt studies of archaeological fish remains while, on the other hand, simultaneous improvement in field recovery techniques through generalised sieving favoured the collection of representative samples. Nonetheless, the analysis of fish bones was hampered by lack of good reference collections needed for identification.

The first samples of fish remains from Natufian (Late Epipaleolithic) deposits were published a few years later. They originated from sites in different settings (Fig. 1): Hatoula, in the western foothills of the Judean hills (Lernau, 1985; Lernau and Lernau, 1991), el-Wad Terrace, in the foothills of Mount Carmel (Valla et al., 1986) and Eynan/Ain Mallaha (Late Natufian occupations), in the Hula Basin (Desse, 1987).

The finding of rare sea fish remains at Hatoula was of peculiar interest because it implied transport over a distance of about 30 km. At el-Wad and Eynan/Ain Mallaha, fish were from the local environment.

The nature and possible intensification of fishing by the Natufians are yet to be fully understood (Bar-Yosef and Zohar, 2010). Besides the relative importance of fish as a food supply, other issues became apparent such as local versus distant captures as well as the environmental significance of the finds. For example, the recovery of fish bones at Hilazon Tachtit (Cyprinidae and Mugilidae families), in western Galilee (Bar-Yosef and Zohar, 2010), and at Baaz Rockshelter (Cyprinidae and Salmonidae) in Syria, today in desert conditions (Conard et al., 2013; Napierala et al., 2013), opens the possibility either of the presence of suitable waterbodies in the vicinity of the sites or relatively distant captures. Finding of specimens assigned to the *Salmo* genus in the Final Natufian layer at Eynan/Ain Mallaha brings further elements to the discussion of these questions.

2. Material and methods

Renewed excavations at Eynan/Ain Mallaha (1996–2005) by F. R. Valla and H. Khalaily (Valla et al., 2007, 2017) produced tens of thousands of fish bones from the Final Natufian layer dated between 10,730 and 9760 cal BC, 2σ (Intcal13; Reimer et al., 2013) (Table 1).

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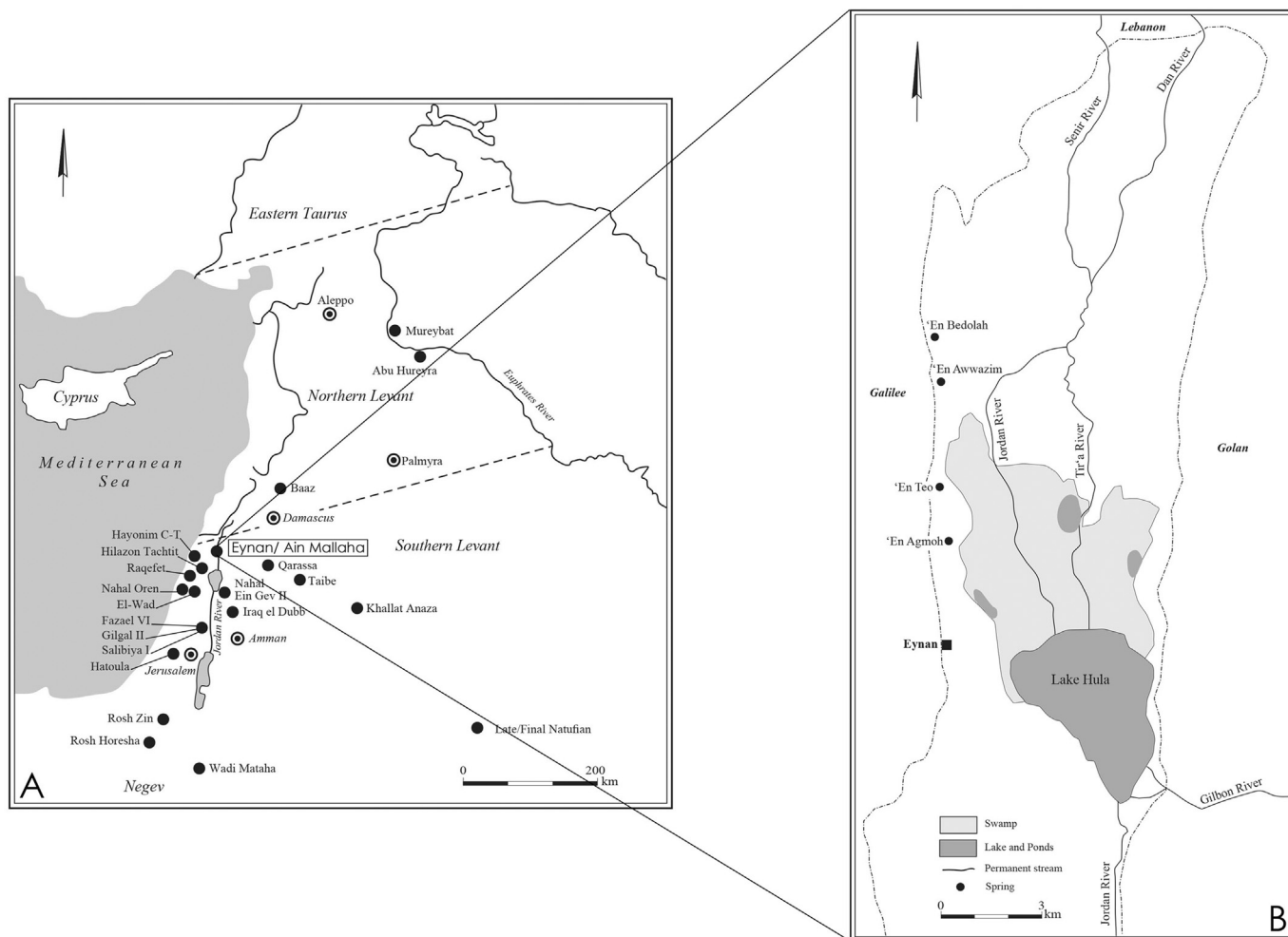


Fig. 1. A: Map of the distribution of Late/Final Natufian sites in the Levant (CAD A. Bridault and H. Khalaily); B: Hydrographic map of the Hula Basin (after Dimentman et al., 1992, modified, CAD H. Khalaily).

Table 1

List of AMS dates of the level Ib (Final Natufian) at Eynan/Ain Mallaha. The results were calibrated based on Intcal13 ¹⁴C calibration data set and calculated by calib.7.0.4 program (Reimer et al., 2013).

Ref. samples	level/struct	ref Lab.	Age BP	sd	cal BC 1 σ (IntCal13)	cal BC 2 σ (IntCal13)
EM05 K95b, 1031-2	Ib/st 230	GifA 70,013	10,200	50	10,061–9858	10,157–9760
EM97 R97 6165	Ib2/st 215	GifA 99,332	10,530	100	10,705–10,433; 10,319–10,294	10,742–10,170
EM99 R98c 7657	Ib2/st 228	GifA 100,400	10,540	90	10,696–10,448	10,743–10,272; 10,268–10,196
	Average Ib2				10,639–10,460	10,732–10,285

The site, in the Hula Basin of the Upper Jordan Valley, is known for its rich Natufian deposits which span most of the duration of this cultural entity (Ashkenazi, 2013; Bouchud, 1987; Bridault et al., 2008; Perrot, 1960, 1966, 1968; Weissbrod et al., 2017). In the recent excavations, the Final Natufian occupation is *circa* 50 cm thick, over a surface of about 120 m² and includes five distinctive stone constructions distributed into two temporally separated building phases (Valla et al., 2007, 2017). A large fireplace from the basal building phase, structure 228 (Fig. 2), dated to 10,740–10,200 cal BC, 2 σ (Valladas and Kaltneker in Valla et al., 2007) (Table 1), was chosen for a pilot study of the fish, due to the abundance of remains in its fill (Valla et al., 2013).

During excavations, systematic wet sieving on 1–2 mm meshes was carried out to collect the fine fraction of the archaeological material. A thorough sorting of the fish remains from fireplace 228 was done by the first author under a stereomicroscope (10–40 \times), guaranteeing exhaustive recovery of the material. Identification was carried out mainly

using reference collections: National Natural History Collections (Hebrew University of Jerusalem, Israel), Dr. I. Zohar's personal collection, and Comparative Anatomy Lab collections of ONIRIS (Nantes Atlantic College of Veterinary Medicine, Food Science and Engineering, France).

3. Results

3.1. Taxonomic composition of the studied sample

Only a part of the fish bones from structure 228 has been studied to date. Already 7475 remains have been analysed and many more are expected to undergo analysis. Out of the analysed sample, 3471 were identified, at least to the family level (Table 2). Most of them were assigned to Cichlidae (Number of Identified Specimens, NISP: 2292) and to Cyprinidae (NISP: 1173). Head bones (mainly from the



Fig. 2. General view of fireplace 228 in wall 225. Photo F. Valla.

Table 2

Taxonomic composition of the studied sample from structure 228 (Number of Identified Specimens - NISP).

Structure 228	NISP	% NISP	%
Cyprinidae	1173	33.8	
Cichlidae	2292	66.0	
Salmoninae	6	0.17	
Identified	3471	100	46.4
Unidentified	4004		53.6
Total	7475		100

splanchnocranium) and vertebrae of both taxa were recovered. Among the Cichlids, lower pharyngeal bone morphology demonstrates the presence of *Coptodon zilli*. The Cyprinidae family is more diverse with four species identified through their pharyngeal bones: *Luciobarbus longiceps*, *Capoeta damascina*, *Garra rufa* and the endemic extinct fish *Mirogrex hulensis*, out of the eight species known from Lake Hula (Zohar et al., 2014).

The most surprising find was that of six vertebrae exhibiting the lateral alveolar pattern characteristic of the Salmoninae sub-family, which includes Atlantic salmon *Salmo salar* and trout *Salmo trutta* complex (Fig. 3), since no representative of this sub-family is nowadays naturally present in the Near East. A scan of the material retrieved from other sectors of the site yield an additional 91 *Salmo* vertebrae, and more are expected. The 97 *Salmo* vertebrae so far identified form the largest record of this taxon in the prehistoric Levant. They nonetheless represent only a small fraction of the total number of fish remains at the site: in structure 228, *Salmo* sp. accounts for only 0.17% of the NISP sampled so far.

The sample of *Salmo* vertebrae consists of 41 thoracic (or precaudal) vertebrae and 50 caudal vertebrae based on the presence or absence of haemal spines, as well as 6 unidentified vertebrae (Table 3).

3.2. The *Salmo* sub-sample

The Salmonidae are distributed throughout the Holarctic region (Bruslé and Quignard, 2013; Sanford, 2000). This family includes three sub-families, Salmoninae, Coregoninae and Thymallinae, easily identified based on vertebrae morphology (e.g. Le Gall, 1984). Within the Salmoninae, the distinction between Atlantic salmon *Salmo salar* and brown trout *Salmo trutta* is more difficult since both species are osteologically very close, with a high intra-specific variability (Desse-Berset, 1994; Guillaud, 2014; Guillaud et al., 2016; Le Gall, 1984). In order to try and differentiate between both species, we used X-ray pictures of the caudal vertebrae as recommended by Desse and Desse (1976) and Desse-Berset (1994). For the brown trout, the vertebral centrum shows a straight “X” pattern, whereas for the Atlantic salmon, it is in the shape

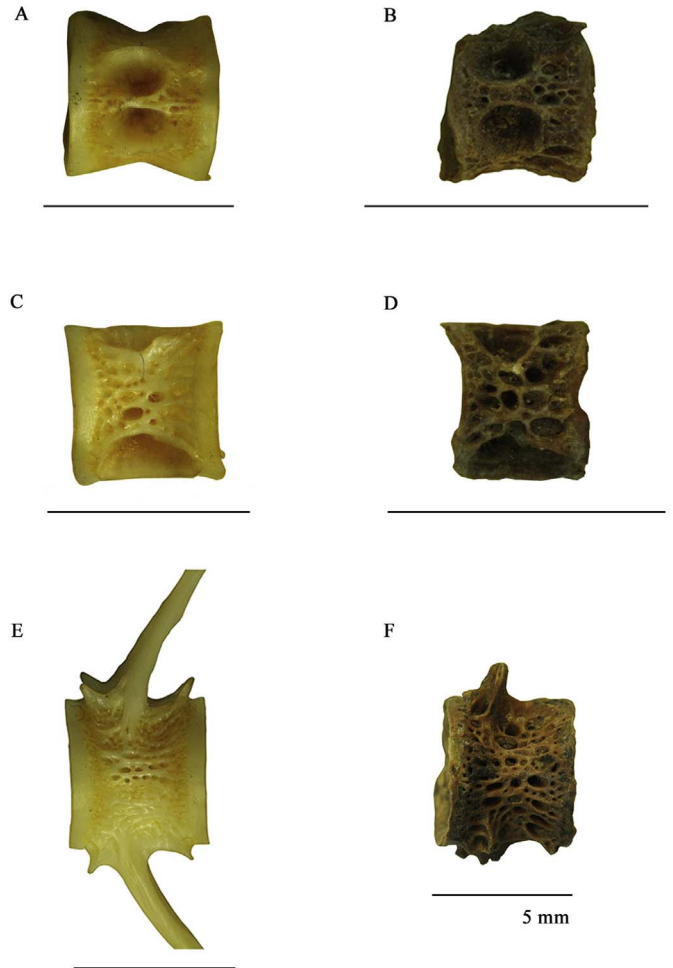


Fig. 3. Thoracic (dorsal and lateral view) and caudal vertebrae (lateral view) of A, C, E: *Salmo trutta* from the reference collection of the Comparative Anatomy Lab (ONIRIS), and for B, D, F: *Salmo* cf. *trutta* from Eynan/Ain Mallaha (thoracic vertebrae: EM96, H94c, 5576; caudal vertebrae: EM97, R97, 5843). The lateral alveolar pattern characteristic of the family is highly variable inside the different trout forms. Photos: A. Borvon.

Table 3

Distribution of the *Salmo* vertebrae in the Final Natufian level at Eynan/Ain Mallaha.

Salmoninae	St. 228	outside St. 228	Total
Thoracic vertebrae	1	40	41
Caudal vertebrae	5	45	50
Vertebrae of indeterminate rank		6	6
Total	6	91	97

of two opposite curved “Y”s (Fig. 4). Eight caudal vertebrae from Eynan/Ain Mallaha were successfully radiographed, and it is clear that their pattern is closer to trout than to salmon (Fig. 4).

4. Discussion and conclusion

Almost 7500 fish remains were studied from a sample originating from structure 228, Final Natufian Eynan/Ain Mallaha. This is the largest sample for the Natufian period for which available publications describe rarely more than a hundred items and usually even no more than a few dozen (Bar-Yosef and Zohar, 2010; Grosman et al., 2016; Van Neer et al., 2005). Cyprinidae and Cichlidae were expected in the context of Eynan/Ain Mallaha and they had already been previously reported (Desse, 1987; Zohar in Valla et al., 2007: 307).

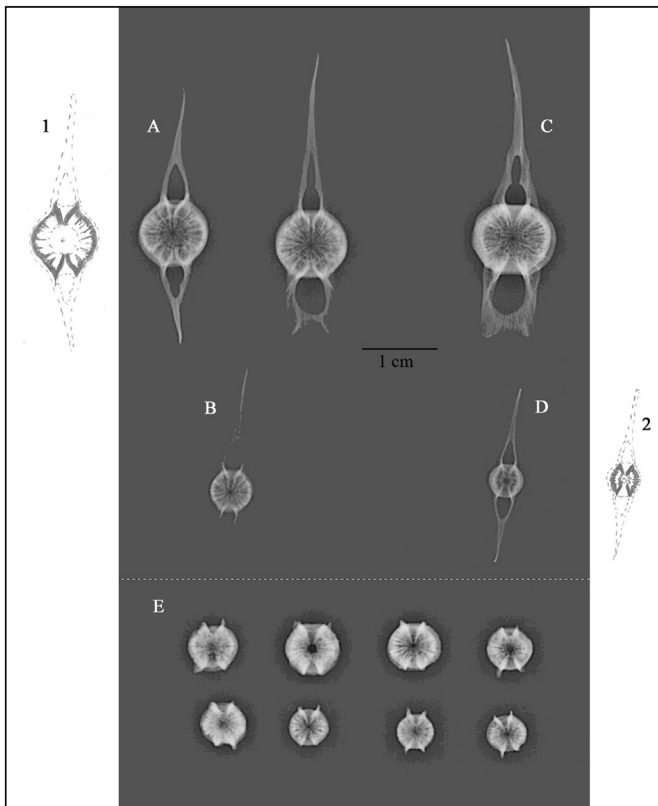


Fig. 4. Caudal vertebrae X-ray (X-ray picture: ONIRIS, Nantes, France) of A, B: Salmon *Salmo salar*, C, D: Trout *S. trutta* from the reference collection of the Comparative Anatomy Lab (ONIRIS), E: *Salmo cf. trutta* from Eynan/Ain Mallaha (from right to left and from top to bottom: EM97, O98d, 5834; EM96, O81, 5178; EM97, Q98, 6105; EM98, S95, 6502; EM97, P97, 5989; EM97, O98b, 6250; EM98, S95d, 7018; EM97, O99, 6251). *Salmo* caudal vertebrae drawing, after the X-ray picture, 1: Atlantic salmon *S. salar*, 2: Brown trout *S. trutta*.

4.1. Anthropogenic origin of the *Salmo* bones

The fish bones at the site were widely scattered and found mixed with the rest of the archaeofaunal remains - mostly food waste, suggesting that fish were part of the diet of the Natufian people. This is reinforced by the finding of a few possible fishing implements, mainly hooks and net weights at the site (Le Dosseur in Valla et al., 2004, Valla et al., 1998). No cut marks have been noted on *Salmo* vertebrae so far, but they are not expected on small-sized fish remains (Wheeler and Jones, 2009; Willis et al., 2008) and burning was observed on five *Salmo* vertebrae. Nonetheless, this may not reflect cooking activities, as bone can be unintentionally burnt when discarded at or near a fireplace (Asmussen, 2009).

No evidence of digestion, which could have raised the question of consumption by predator species, was noted (Jones, 1984, 1986; Nicholson, 1993, 2000). The fact that only vertebrae were recovered is probably related to the fragility of the head skeleton in the Salmoninae group, as it is seldom recovered from archaeological deposits (Butler and Chatters, 1994; Lubinski, 1996; Morales and Rosello, 1998). All in all, trout were apparently very unusual in people's meals.

4.2. Which *Salmo* species?

In addition to the osteological criteria, the attribution of the vertebrae to the Atlantic salmon can probably be excluded when considering its indigenous range: mainly in the northern Atlantic Ocean (Klemetsen et al., 2003; Porcher and Baglinière, 2011) (Fig. 5). It has been suggested, on the basis of archaeological finds, that salmon populations were resident in the western Mediterranean Sea from the Last

Glacial Maximum (21 ka cal BP) to the early Holocene (Guillaud, 2014; Kettle et al., 2008, 2011; Le Gall, 2008, 2010). To date, this is an open issue (Guillaud et al., 2016).

At Eynan/Ain Mallaha, a member of the trout group is a more likely candidate when considering their geographical distribution (Fig. 5). In the Near East, trout is recorded in the Upper Euphrates and Tigris Basins of southern Turkey (Baglinière, 1991; Bruslé and Quignard, 2013; Ombredane et al., 2011).

According to DNA studies, most European trout seem to belong to a dominant species, *Salmo trutta*. This species is constituted of an aggregate of interbreeding forms, each centred on a particular geographical setting but with considerable overlap (Bernatchez, 2001; Cortey et al., 2004, 2009; Ombredane et al., 2011). A few forms, very located, are still given a status of species, like *Salmo platycephalus* in southern Turkey and *Salmo tigridis* in the Tigris Basin, for the Near Eastern morphotypes (Sušnik et al., 2004, 2006; Turan et al., 2011). However no osteological criteria are available to differentiate them from the brown trout.

4.3. Ecology

As it is highly adaptable, *Salmo trutta* is a ubiquitous species that can be found in seas, lakes and rivers (Ombredane et al., 2011). However, sea trout are a migratory variety, whereas lake and river trout do not tolerate salt water. Their distribution is also limited by some physical parameters, mainly water temperature and oxygenation.

Trout favour well oxygenated water between 0° and 20 °C, with optimal temperatures between 5° and 17 °C, but they do not survive water above 22°-26 °C (Bruslé and Quignard, 2013; Elliott, 1982; Jonsson and Jonsson, 2009; Ombredane et al., 2011). In reproduction areas, water should not exceed 10 °C. Thus, in rivers, spawning places are usually upstream.

4.4. Did physical conditions in the vicinity of Eynan/Ain Mallaha 12,000 years ago match these requirements?

To this day, the Hula depression is rich in fresh water (Fig. 1B). Many springs flow into its margins at the foot of the Galilee and Golan Heights, permanent streams run from the Anti-Lebanon Mountains, and the remnants of the lake occupy its lowest place. In recent past, the spring of Ain Mallaha presented a stenothermic temperature of 21–22 °C (Dimentman et al., 1992: 89). The Lake Hula was 3 to 4 m deep, with temperatures varying according to time of the day and seasons between 5° and 37 °C. Its oxygenation was poorly documented (Dimentman et al., 1992: 19, 26–31). Its catchment extended north in adjacent mountains including Mount Hermon (Dimentman et al., 1992: 23). In such areas, as well as in the nearby upper Galilee, trout could have found suitable habitats even under modern conditions (Gravel, 1931). This is also confirmed by the modern breeding of the rainbow trout *Oncorhynchus mykiss*, in such streams as Dan River (Fig. 1B) with temperatures today of 15 °C all year round (Golani comm. pers. 2016).

The situation in the area during the Final Natufian can be reconstructed through archaeofaunal and palaeobotanical analyses. Some studies have pointed out the relative stability of environment (Ashkenazi, 2013; Biton et al., 2016), while other support a somewhat cooler climate in comparison to nowadays (Bar-Matthews et al., 1997, 1999, Hartman et al., 2016, Mienis in Valla et al., 2007). Generally speaking, as Tchernov (1997: 221) emphasised: “no real dry periods existed during the Late Pleistocene in the Southern Levant but [...] during this period the Southern Asian arid belt was always much wetter supporting a rich diversity of Palearctic species [...]”. In the Hula, the wide variety of taxa recovered in layer Ib at Eynan/Ain Mallaha supports this view. Accordingly, even if it is doubtful that trout found optimal conditions in the immediate vicinity of the site, it is likely that they could live locally, at least in nearby hilly areas.

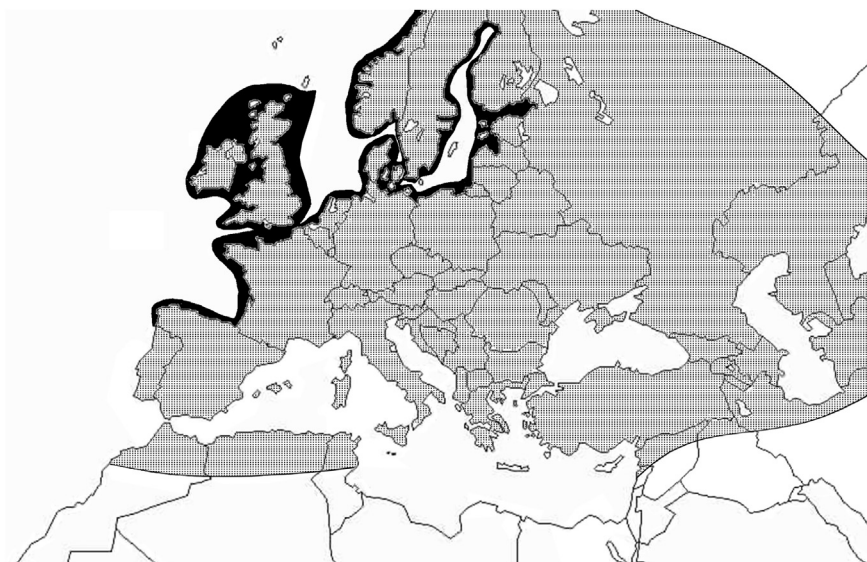


Fig. 5. Current spatial distribution of Atlantic salmon *Salmo salar* in black and brown trout *Salmo trutta* complex in grey (map after Baglinière, 1991, Bernatchez, 2001, Porcher and Baglinière, 2011, modified, CAD C. Picard). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

4.5. Conclusion

All in all, it is likely that trout could have lived in the vicinity of Eynan/Ain Mallaha at the end of the Pleistocene. An alternative possibility, namely introduction by transport from further north, cannot be totally excluded in a context where trade of Cappadocian obsidian is well attested (Khalaily and Valla, 2013). Nevertheless, the most parsimonious hypothesis is their local presence, at least in the cooler rivers flowing into the Hula Valley from the north.

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