THE CONTRIBUTION OF SPRINGTIME AND AUTUMN GLASS EELS (ANGUILLA ANGUILLA) TO STOCK: RESULTS BASED ON OTOLITH MORPHOMETRY.

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ABSTRACT

A double mark, called a transition ring, or elver mark, was identifiable in light microscopy on otoliths of young yellow eels. In the Vilaine watershed, the radius of this mark decreased from 178 µm in yellow eels corresponding to glass eels arriving in autumn 1997 to 163 µm in yellow eels arriving in spring 1998. The mean transition ring radius of the freshwater eel population in the Vilaine river had an intermediate value between spring and autumn recruits. This implies that it consisted of a mixture of spring and autumn recruits. In the Vilaine estuary and the Frémur populations, the mean radius of the transition rings was close to the autumn one. The springtime recruits formed 68% of freshwater and 15% of estuarine population in the Vilaine for the 1998 cohort. This result was in sharp contrast with the available assessments of recruitment, which both in estuary and in the fluvial part of the watershed, were dominated by spring recruits. This contrast is possibly the consequence of density-dependent mortality, which would have been particularly important in springtime for glass eels whose migration was inhibited by the dam.

Key-words: Anguilla anguilla, eel, mortality, transition ring, season, estuary.

CONTRIBUTION DES RECRUTEMENTS PRINTANIERS ET AUTOMNAUX DE CIVELLES (ANGUILLA ANGUILLA) : RESULTATS BASÉS SUR LA MORPHOMÉTRIE DE L’OTOLITHE.

RÉSUMÉ

Une double marque, nommée anneau de transition est identifiable en microscopie optique sur les otolithes des jeunes anguilles. Sur le bassin versant de la Vilaine, le rayon de cette marque passe de 178 µm chez les anguillettes correspondant à des civelles arrivées à l’automne 1997, à 163 µm chez des anguillettes correspondant à des recrues du printemps 1998. Le rayon moyen de la marque de transition de la population d’anguille fluviale de la Vilaine est intermédiaire entre celui des recrues printanières et automnales. Ceci implique que les populations d’anguille fluviale de Vilaine résultent d’un mélange de recrues printanières et automnales. Mais pour les populations de l’estuaire de Vilaine et du Frémur le rayon de transition est proche de celui des recrues automnales. La proportions de recrues printanières s’établit à 68% en eau douce et 15% en estuaire pour la cohorte 1998 en Vilaine. Les recrutements fluviaux et

Article available at http://www.kmae-journal.org or http://dx.doi.org/10.1051/kmae:2003034
estuariens de civelles correspondant à l’échappement à la pêche sont au contraire dominés par les civelles printanières qui forment l’essentiel de l’effectif. Cette opposition pourrait provenir d’une mortalité densité-dépendante, particulièrement importante au printemps pour les civelles bloquées au pied du barrage.

Mots-clés : Anguilla anguilla, anguilles, mortalité, anneau de transition, saison, estuaire.

INTRODUCTION

The decline in glass eel (Anguilla anguilla) recruitment on the Atlantic coasts (MORIARTY and DEKKER, 1997) may result in a change in yellow eel population structure, density and geographical extent, at the edge of the distribution area. However, these changes may not occur in situations where density dependent factors moderate the effect of a fall in recruitment (SVEDÄNG, 1999; KNIGHTS et al., 2001). Atlantic facing estuaries form a particular context because they receive, as a rough estimate, 76% of the total recruitment but only account for 7% of the distribution area in Europe (DEKKER, 2000). In these locations, the short duration of the estuarine migration implies that only juvenile yellow eels colonize the watershed, and that glass eels remain for a while in, or close to, the estuary. During this period, density dependent regulations will affect both mortality and migration. Mortality of both glass and juvenile eels is determined by the difference between carrying capacity and recruitment. Density dependent regulations will also play a role in the colonisation process as population pressure in the lower reaches of the catchments would enhance upstream migration (MORIARTY, 1986; LOBÓN-CERVIÁ et al., 1995). The probability for a glass eel of surviving may vary depending on its energy reserves (CHARLON and BLANC, 1982; BOËTIUS and BOËTIUS, 1989; GUÉRAULT et al., 1995; DE CASAMAJOR et al., 2001a). Glass eels entering the estuary from August to November have the best condition factor and are present at a lower density during glass eel stage than glass eels arriving in winter or springtime. Therefore, they are supposed to have a better survival rate.

The aim of this paper is to analyse the respective contribution to the watershed subunit of stock of early autumn recruits compared to spring recruitment in the Vilaine watershed. In this river, almost no glass eel escapes capture from the fishery during the fishing season from mid-December to April (BRIAND et al., In press). Recruitment is therefore limited to the small quantities of autumn glass eel arriving before and at the beginning of the fishing season, in November, when fishing is only conducted by a few boats. Most of the recruits consist of spring glass eels arriving after the closure of the fishery.

A method for differentiating spring from autumn recruits at the adult stage based on otolith measurements is described in this paper. For all eel species, the transition ring (or elver mark) is formed on glass eel otoliths at the end of marine life (MICHAUD et al., 1988; LECOMTE-FINIGER, 1992; TZENG and TSAI, 1994; KAWAKAMI et al., 1998). This ring does not strictly correspond to the entry to freshwater, as it may already be observed in glass eels collected far from the coasts (LECOMTE-FINIGER et al., 1993; DE CASAMAJOR et al., 2001b); or deposited experimentally in full salinity water (CIERI and MC CLEAVE, 2001). In 1990, GUÉRAULT et al. (1992) described a seasonal decrease of the radius of the glass eel otolith, measured from the centre to the transition ring. This result suggests that it could be possible to distinguish the respective contributions of spring and autumn recruits to the whole population.

The objectives of this work are to show that there has been a seasonal decrease of the dimensions of the transition ring from autumn 1997 to spring 1998. We will identify two groups of yellow eels, one recruited in autumn and one in springtime. The
distributions of the transition ring radius of three populations of yellow eels - collected on the freshwater parts of the Frémur and the Vilaine and in the Vilaine estuary - will then be compared with the spring and autumn groups previously described. Finally we will calculate the contribution of spring and autumn recruits to the 1997 and 1998 cohorts in the Vilaine. This contribution will be compared with the recruitments recorded at the glass eel stage and the possible consequences on glass eel mortality will be analysed.

MATERIALS ET METHODS

Description of the study areas

The Vilaine watershed is located on the southern coast of Brittany, in the North-West of France and covers 10 400 km². The eel population is separated into two distinct compartments by the Arzal dam, which delimitates the estuary, 12 kilometres from the river mouth (MOUNAIX, 1992). Glass eels concentrate at the bottom of the dam where the professional fishery applies heavy fishing pressure. Upstream from the dam, the Vilaine forms a 200 m wide river and separates into two tributaries (Figure 1). An eel-trapping ladder is located on the first dam of each stream, 45 and 80 kilometers from the estuary.

The Frémur watershed is a short coastal stream (60 km²) located on the north coast of Brittany (Figure 1). It is obstructed by several dams, but the main one is equipped with both an eel lift and a trapping system (Bois Joli, 5 km from the estuary). The fishing pressure in the short estuary is very limited as there is no professional fishery and angling mainly focuses on game fish (FEUNTEUN et al., 1998).

Sample collection

We sampled 217 yellow eels in the Vilaine estuary. They were collected in 1998 and 1999 on the estuarine eel ladder, and in 1999 in the estuary from a commercial eel
pot fisherman. We also collected 549 eels in the freshwater part of the Vilaine watershed in 1998 and 1999. Most eels were trapped on the two eel ladders located in the watershed but a small sample was added from an electrofishing operation in 1999. We also collected 120 eels by electrofishing from the freshwater part of the Frémur watershed in 2000 (Table 1). A total of 886 eels was therefore used for otolith extraction.

Table 1
Collection date and capture locations of eels used for otolith measurements.

<table>
<thead>
<tr>
<th>Place and date</th>
<th>N</th>
<th>Sample (size range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vilaine estuary</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Estuarine trapping ladder, April-May 1998</td>
<td>47</td>
<td>( random) 70-220 mm</td>
</tr>
<tr>
<td>Estuarine trapping ladder, Oct-Nov 1999</td>
<td>96</td>
<td>Size graded (all sizes)</td>
</tr>
<tr>
<td>Eel pot (fisherman) Oct-Nov 1999</td>
<td>74</td>
<td>Size graded (all sizes)</td>
</tr>
<tr>
<td>Vilaine Freshwater</td>
<td></td>
<td></td>
</tr>
<tr>
<td>River Claie, 80 km, electrofishing, Nov 1999</td>
<td>30</td>
<td>Random (all sizes)</td>
</tr>
<tr>
<td>La Potinais, 45 km, trapping ladder, June-Sept 1998</td>
<td>123</td>
<td>Random &lt;120 mm (66-111 mm)</td>
</tr>
<tr>
<td>La Potinais, 45 km, trapping ladder, May-Sept 1999</td>
<td>285</td>
<td>Random &lt;150 mm (65-139 mm)</td>
</tr>
<tr>
<td>Malon, 70 km, trapping ladder, 1999, June-Aug 1999</td>
<td>111</td>
<td>Random &lt;150 mm (82-150 mm)</td>
</tr>
<tr>
<td>Frémur freshwater</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Electrofishing 2000</td>
<td>120</td>
<td>Size graded (70-380)</td>
</tr>
</tbody>
</table>

Otolith processing

Otoliths were stored dry. They were observed using a binocular microscope (Olympus, x1 to x63) under reflected light. To enhance contrast, they were immersed in 70 % ethanol. Otoliths of eels larger than 150 mm were embedded in resin (methacrylate, Dentsply) and mechanically ground (Dremel grounder, 600 grit emery disk) down to the sagittal plane before observation (MOUNAIX, 1992). Image analysis was used to identify and measure the radii of growth structures on otoliths. The inner translucent zone delimited the nucleus whose border corresponded to the transition ring (LECOMTE-FINIGER et al. 1992; GUÉRAULT et al. 1992). Outward from the nucleus, the other measured marks (annuli=winter rings), have been described by PANFILI and XIMÉNÉS (1994) as the contrasting limit between translucent and opaque zones. They corresponded to fine translucent lines located at the beginning of each opaque zone (Figure 2).

Seasonal group identification

Spring eels were defined as glass eels recruited in springtime after the end of the fishing season (i.e. March to July). 1998 spring recruitment was marked with tetracycline hydrochloride (TCHC, method in ALCOBENDAS et al. 1991). A total of 210 kg of glass eels caught on the estuarine ladder in May and June 1998 (30 % of the spring recruitment) were marked. They were then released in freshwater above the ladder and next recaptured on the freshwater trapping ladders located upstream in the watershed. The tetracycline hydrochloride mark (ALCOBENDAS et al., 1991) was detected on otoliths using an epifluorescent light microscope (LEITZ Laborlux S, objective X10, eyepiece x40). When exposed to U.V. light (excitation filter 450-490 nm) the fluorochrome appeared as a bright yellow circular line in the otolith of marked eels (stop filter: 515 nm).
Juvenile eels were also collected in May and June 1998 on the Arzal estuarine eel ladder with the purpose of identifying Autumn 1997 recruits within the sample, on the basis of body size and otolith characteristics.

Contribution of the seasonal groups to the population

We measured the radii of the transition rings in the Vilaine freshwater, the Vilaine estuary and the Frémur (freshwater) samples. Their distributions were compared with one another and with spring and autumn recruits by Kolmogorov Smirnov non-parametric tests, because of the lack of homoscedasticity of data distributions. Next, we calculated the respective proportions of spring and autumn recruits within the estuarine and freshwater populations of the Vilaine. The calculation was based on transition ring radii obtained in autumn 1997 and spring 1998 samples. The proportion (p) of spring recruits was calculated by the following relationship: $M = p m_1 + (1-p)m_2$, with $M$= mean transition ring radius of freshwater or estuarine population, $m_1$ = mean transition ring radius of spring population, $m_2$=mean transition ring radius of autumn population.

In order to analyse the effects of cohort and location on transition ring variation, an analysis of variance (GLM SYSTAT) was performed on 853 transition ring radius values. Five cohorts (1995 to 1999) calculated from age and year of capture were selected. Data were also split into three sampling locations, Vilaine estuary, Vilaine freshwater and Frémur. The following model was used:

$$\text{transition ring} = \text{constant} + \text{location} + \text{cohort} + \text{cohort} \times \text{location}.$$
RESULTS

Sample collection and otolith processing

On 886 eels, a total of 882 otolith pairs were processed, including 215 from the Vilaine estuary, 549 from freshwater bodies of the Vilaine river and 118 from the Frémur watershed system. Otolith observations provided measurements of 853 transition rings, 29 were not interpretable due to failure in otolith preparation.

Figure 3
Size (mm) and age distributions of glass eels and juvenile eels migrating at the Arzal trap. Size frequencies were calculated on a sample of at least 200 juvenile eels and extrapolated to the whole monthly catch. Glass eel migration in spring 1998 amounted to more than two million eels and is only indicated on the graph. The arrows correspond to the expected size of autumn recruits. Bottom right, size frequency of eels for which the winter 1997 translucent slow growth ring is not visible in spring 1998 (assumed autumn recruits).

Figure 3
Histogramme de fréquences de tailles et décomposition en âge des civelles et anguillettes en migration sur la passe d’Arzal. Les fréquences de tailles sont calculées sur la base d’un échantillon d’au moins 200 anguilles et extrapolées à l’ensemble de la capture mensuelle. La migration de civelles au printemps 1998 représentait plus de deux millions d’individus et est juste indiquée sur le graphique. Les flèches représentent la taille supposée des recrues automnales. En bas à droite, fréquence de tailles des anguilles dont l’anneau translucide de l’hiver 97 n’est pas visible au printemps 98 (recrutement automnal supposé).
Seasonal group identification

Only 38 yellow eels were observed with one tetracycline mark and thus identified as 1998 spring recruits. The identification of 39 eels as autumn eels was based on body size, developmental stage and otolith criteria. In May and June on the Arzal eel ladder, we could easily distinguish spring recruits still at the glass eel stage from fully pigmented juvenile eels (stage VII ELIE et al., 1982). During the fishing season in the Vilaine the fishing pressure is very high. As a consequence, only the "young" i.e. early pigment stages are found viz. VB and VIA0. Old stages start to appear only after the end of the fishing season in April and May. In spring samples, the young yellow eel advanced stage identified them as being a different group from spring 1998 recruits and they were very likely recruited the previous year.

Between the young yellow eels, otolith criteria were then used to differentiate between autumn recruits and spring recruits of the previous year. Some eels had otoliths on which the marginal deposition of the opaque zone had not started, so the identification of the translucent winter ring was not possible. These eels were small with sizes ranging from 70 to 130 mm. The eels with an opaque margin on otolith were larger with size ranging from 110 to 200 mm. We assume that spring deposition of the opaque zone occurs later in eels arrived in autumn because of a slow growth resumption. Therefore, juvenile eels without opaque margin on otoliths were identified as eels recruited as glass eels in autumn.

Size and age distributions of monthly catches of migrating eels at Arzal dam are in accordance with those observations (Figure 3). The glass eels recruited in autumn grew slowly during winter to reach in May-June the size of small juvenile eels on whose otoliths the opaque margin had not yet been deposited (bottom right-hand histogram, Figure 3).

This assumption was further confirmed by the measurement of the radius of the transition ring, the largest rings being identified in the autumn group. The radius of the transition ring was significantly lower in spring recruits than in autumn recruits (means 163.4 and 178.3 µm respectively, Kolmogorov Smirnov, P<0.001). This difference between autumn and spring recruits indicated that there was a seasonal variation in the transition ring and that this variation was still measurable at the yellow eel stage (Figure 4).

Contribution of the seasonal groups to the population

In samples coming from the Vilaine watershed, the transition ring radius distributions were different (KS p<0.05) between freshwater and the estuary. Their means (169.9 and 176.6 µm, respectively) were intermediate between typical spring and autumn values. In the Frémur, the radii of transition rings were the largest (mean 178.0 µm); and their distribution was different from both Vilaine freshwater (KS p<0.01) and Vilaine estuary (KS p<0.05) (Table 2).

Within estuarine samples, the transition ring distribution of eels collected on the eel ladder (mean 176.2 µm) and in the estuary (mean 177.6 µm) were not different (p>0.5 KS) (Table 2).

The proportions of spring recruits (p) were calculated from transition ring radii in the 1998 cohort as being p=68 % for freshwater and p=15 % for estuary.

In the GLM analysis, the crossed cohort*group effect was not significant. In the Vilaine basin, freshwater and estuarine samples were different (p<0.001, Bonferroni) (Figure 5A). There was however no difference between cohorts (Tukey p>0.5) (Figure 5B). Pairwise comparisons of cohort*location crossed effects were not significant except for the difference between freshwater 1998 and estuary 1996 (Tukey p<0,05) (Tables 3, Figure 5C).
Table 2
Measurements of transition ring radius of otoliths collected in the Vilaine in estuary (estu98-99), river (fresh98-99), in the groups corresponding to early glass eels (Fall97) and late (Spring98) arrivals, and on the Frémur (Frémur00). Kolmogorov Smirnov test statistics for between group comparison, *** p<0.001, ** p<0.01, * P<0.05, NS non significant.

<table>
<thead>
<tr>
<th></th>
<th>N</th>
<th>Mean (µm)</th>
<th>Min (µm)</th>
<th>Max (µm)</th>
<th>SD</th>
<th>Spring</th>
<th>Fresh</th>
<th>Estu</th>
<th>Fall</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spring 98</td>
<td>38</td>
<td>163.4</td>
<td>117</td>
<td>190</td>
<td>14.8</td>
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<tr>
<td>Fresh 98-99</td>
<td>512</td>
<td>169.9</td>
<td>122</td>
<td>230</td>
<td>16.5</td>
<td>*</td>
<td>.</td>
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<td>.</td>
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<tr>
<td>Estu 98-99</td>
<td>177</td>
<td>176.6</td>
<td>119</td>
<td>247</td>
<td>20.9</td>
<td>***</td>
<td>*</td>
<td>.</td>
<td>.</td>
</tr>
<tr>
<td>Fall 97</td>
<td>39</td>
<td>178.3</td>
<td>140</td>
<td>230</td>
<td>19.2</td>
<td>***</td>
<td>**</td>
<td>NS</td>
<td>.</td>
</tr>
<tr>
<td>Frémur 00</td>
<td>116</td>
<td>178.0</td>
<td>149</td>
<td>213</td>
<td>13.8</td>
<td>***</td>
<td>***</td>
<td>*</td>
<td>NS</td>
</tr>
</tbody>
</table>

Figure 4
Distributions of the transition mark radius of 38 yellow eels corresponding to spring recruits and 39 yellow eel corresponding to autumn recruits.

Figure 4
Distributions des rayons des marques de transition sur les otolithes de 38 d’anguilles correspondant à des recrues printanières et 39 anguilles correspondant à des recrues d’automne.
The first step in this work was to show that there was a seasonal variation of transition ring radius in glass eel otoliths, and that this variation could still be observed in young yellow eels. The tetracycline mark provided an unambiguous identification of eels belonging to the spring recruitment. The identification of autumn recruits was less easy. Within yellow eels collected in the springtime on the eel ladder, some were very small, but could not be confused with spring recruits, which were still at the glass eel stage. Indeed, spring recruits were freshly arrived in estuary, i.e. in April, because there is almost no escape from the fishery before its closure (BRIAND et al., In press). Among yellow eels, the boundary between autumn recruits (age 0) and spring recruits arrived the year before (age 1) was more difficult to establish and was based on otolith shape criteria.

To resume with, in spring, glass eels freshly arrived had otoliths with little marginal growth beyond the transition ring. Autumn recruits had a larger margin. In both groups, the deposition of the opaque marginal zone corresponding to the spring period of rapid growth was not sufficient to distinguish the translucent ring at the margin of the otolith. In opposition, eels from the previous springtime had a large marginal opaque zone deposition and displayed a translucent ring.

The second step was to test whether the differences between autumn and spring glass eels were stable in time. Indeed, the seasonal variations of the transition ring radius are not necessarily stable between years according to GUÉRAULT et al. (1995) and DE CASAMAJOR et al. (2001b). The outcome from these results was that a yearly stability of the seasonal trend seemed doubtful, and that analysis should be limited to the 1997-1998 cohort for which we had obtained values of the spring and autumn radius.

The long-term variation of otolith characteristics has also to be taken into account. Our results show that there has been a decrease in transition ring radius in the Vilaine since 1990 (GUÉRAULT et al., 1992; GUÉRAULT et al., 1995). The same trend was observed on the Adour (DE CASAMAJOR, Pers. comm, DE CASAMAJOR et al. 2001b). These variations in otolith size corresponded to the general long-term decrease in glass eel size and condition (DEKKER, 1998), which could be due to oceanic events (DÉSAUNAY and GUÉRAULT 1997; DE CASAMAJOR et al. 2001b).

However, these temporal variations were not detectable during our study, as the GLM showed that the temporal effect was not significant (Table 3). The absence of a significant temporal effect may partly be explained by the increasing variance of

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>df</th>
<th>F-ratio</th>
<th>P</th>
</tr>
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<td>Location</td>
<td>2</td>
<td>6.345</td>
<td>0.002</td>
</tr>
<tr>
<td>Cohort</td>
<td>4</td>
<td>0.465</td>
<td>0.761</td>
</tr>
<tr>
<td>Cohort*Location</td>
<td>8</td>
<td>0.239</td>
<td>0.983</td>
</tr>
<tr>
<td>Error</td>
<td>838</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Figure 5

GLM analysis for model transition ring radius=location + cohort + cohort*group. A/ Least mean squares + SE and Bonferroni test results for difference between groups, Vil fresh=Vilaine freshwater, Vil estu=Vilaine estuary, Frémur. Comparison with Vilaine Freshwater, *** p<0.001, * p<0.05. B/ Least mean squares + SE and Tukey test results for difference between cohorts, no significant difference found. C/ Least mean squares + SE and Tukey test results for difference between crossed effect (cohort*group). No significant difference between groups except one case: p<0.05 between Vilaine estuary 1996 and Vilaine freshwater (*).

Analyse GLM du modèle transition ring radius=location+cohort+cohort*group. A/ Ajustement des moindres carrés, + erreur type et test de comparaison de moyenne de Bonferroni pour la différence entre group, Vil fresh=Vilaine fluviale, Vil estu=Vilaine estuaria, Frémur. Comparaison à Vilaine fleuve, *** p<0.001, * p<0.05. B/ Ajustement des moindres carrés + erreur type et test de comparaison de moyenne de Tukey pour la différence entre cohortes (pas de différence significative). C/ Ajustement des moindres carrés + erreur type et test de comparaison de moyenne de Tukey pour la différence entre effets croisés (cohort*group). Une seule différence p<0.05 entre Vilaine estuaria 1996 et Vilaine eau douce 1998 (*)
measurements as the eels grow older and otoliths get larger and more difficult to observe (figure 5B). The remarkable stability in estuarine transition ring radius implies that there was a year-to-year stability in the process coupling seasonal decrease in transition ring size, mortality and recruitment (Figure 5C).

As a consequence, we conclude that the interpretation of the otolith characteristics in young yellow eels, as described in this paper, provides a reliable method for tracing and determining the season of arrival of glass eels in estuaries. Such a result has already been observed in the silver eel (HOLMGREN et al., 1997) and suggests that mortality may smooth seasonal or annual variations in transition ring radius.

The otolith transition ring distributions varied significantly between sites suggesting that the stocks are composed of a variable proportion of spring glass eels (figure 5A). The lowest proportion of autumn eels was found in the freshwater part of the Vilaine, whereas the Vilaine estuarine eels and the Frémur freshwater eels display a large transition ring radius close to the autumn characteristics.

There is a large difference between the freshwater habitats of the Vilaine and the Frémur. So the differences in seasonal groups contributions could be explained firstly by invasion behaviours depending on sites: the Frémur river is much shorter and most sampling sites are more rapidly accessible for autumn eels compared to the Vilaine watershed. The other point is that there is no glass eel fishery in the Frémur River system and no related fishing mortality. A higher proportion of autumn and winter glass eels thus survives and colonises the freshwater habitats. The last point is that a different migration route in the sea between north and south Brittany may lead to a geographical discrepancy in transition ring size.

Within the Vilaine River system, the higher proportion of autumn eels in the estuary than in freshwater can be explained by the Arzal dam, which isolates the freshwater and estuarine populations, and leads to a difference in population dynamics in each compartment. The temporal stability in transition ring radius may be contrasted with the year-to-year variation in recruitment observed on the Arzal eel ladder (BRIAND et al., In press). There may however be a relationship between recruitment and transition ring radius: the smallest transition rings occurred in 1996 both in the Vilaine and in the Frémur freshwater habitats. In both systems this was the year when the dams were equipped with eel ladders. These results suggest that the access to low-density habitats was improved and that this improvement mainly eased spring recruits. In addition, in the Vilaine catchments, the 1998 cohort, which represented the highest fluvial recruitment (700 kg of glass eels), had a slightly smaller transition ring than the 1997 and 1999 cohorts, which had low recruitment levels (70 and 250 kg) (BRIAND et al., In press). This low transition ring value was highlighted by the only significant difference found in pairwise comparisons of location*cohort groups which included the 1998 fluvial cohort (Figure 5C).

The striking outcome from these results is the importance of the proportion of autumn recruits in the Vilaine and the Frémur. On the one hand, calculations from transition ring radius imply that the freshwater yellow eel population consisted of about 32% autumn eels and that this proportion increased to 85% in estuary. On the other hand, focusing on glass eel recruitment, autumn recruits only accounted for 0.2 to 0.6 % of the total fluvial recruitment (BRIAND and FATIN, 2002). The discrepancy between the recruitment estimation and the final population level may indicate: (1) that there was a problem in using transition ring radius, or (2) that the autumn recruitment was not well recorded, or (3) that there has been a large differential mortality between spring and autumn glass eels.
The calculation of the respective contribution of each cohort to the final population has to be considered as a rough calculation because of the small size of the seasonally related samples. However, because they were based on samples of spring and autumn eels collected at the yellow eel stage, the mean transition ring radius can be considered as a valid measurement. Indeed, the eels identified as autumn recruits within the spring ladder sample, displayed a wide range of sizes (80 –120 mm) and a wider range of transition ring radius than the spring sample. These variations were possibly the result of a variable growth, but could also have been the result of a mixing of several waves of glass eels with different transition ring radii and different survival conditions from August to November. The yellow eel sample therefore integrates the final contribution of autumn glass eels to the population.

In the same way, the yellow eels identified as spring recruits corresponded to the marking from May to June of the main run of glass eels recorded on the ladder, and were representative of this run.

In freshwater, the recruitment was carefully monitored on the eel ladder, and the very limited number of glass eels migrating from August to November could not explain the very large autumn eels contribution. However, three other sources of autumn recruits could be found. The first was made of yellow eels migrating through the eel ladder and the vertical slot fishway. Their number varied from 9 540 to 35 320 eels per year from 1996 to 1999 (BRIAND and FATIN, 2002). These yellow eels had a large transition ring radius. Indeed, there was no difference in transition ring size between samples collected on the trap and in the estuary and the otolith shapes were identical, with large and regular spaces between the winter marks. The second source corresponded to the migration at the glass eel stage through the sluice. The annual glass eel weight was evaluated to range from 10 to 30 kg in 1997 and 1999 (unpublished results). This additional recruitment was not large when compared to the eel ladder trapping as a whole, but it outnumbered autumn recruits. The third and probably main source of autumn recruits was the passage through the sluice and dam overflow panels at the yellow eel stage. The size of this recruitment could not be estimated but it was supposed to be large as eels use sluices during their migrations (BARAS et al. 1996; VERDON and DESROCHERS 2000).

In the estuary there was a large uncertainty in recruitment level. From August to November some glass eels were always caught on the eel ladder and this indicates that there was a continuous recruitment in the estuary. In November, at the beginning the fishing season, the fishing pressure was low with only one to ten boats per night, and there was possibly an escapement (BRIAND et al., In press). But as the fishing pressure increases in December, a large survival of November glass eels seems doubtful.

Therefore, in both cases, the size of autumn recruitment was probably very limited when compared to spring recruitment.

As the two previous points fail to explain the large contribution of autumn eels to the freshwater and estuarine population, the third assumption is probably right and a very large mortality of spring glass eels settling in the estuary may explain their small contribution to the stock. This large mortality may be the consequence of a size-dependent mortality, exacerbated by density-dependent factors. Autumn recruits have larger energy reserves than spring eels (BOËTIUS and BOËTIUS, 1989; GUÉRAULT et al., 1995). They also display a better survival and growth rate in rearing experiments than glass eels arriving from January to March (SANTOS and WEBER, 1992). In addition, they arrive a few months before spring recruits in the estuary, and in spring they have already completed their metamorphosis. Size-dependent mortalities are reported in juvenile cod, with a survival of a 0 group released at sea increasing from 2 to 75% when the size at release was increased from 8 to 12 cm (DEKKER, 1998;
KRISTIANSEN et al., 2000). In a review of mortality rates encountered by estuarine glass eels, KNIGHTS et al. (2001) noticed «Natural mortality of glass eel is relatively high and exacerbated by density-dependent effect during the time between their entry into estuaries and the metamorphosis and migration of elvers. Density-dependent Mortality would be exacerbated if migration is delayed in long estuaries or inhibited by barriers, e.g. as in the Severn and Vilaine respectively».

This paper shows that seasonal changes of transition ring radius may provide a tool for investigating eel population dynamics at the glass eel/ young yellow eel stage. Because of overlapping distributions, it will not be possible to differentiate seasonal recruits at the individual level, but results may be obtained at the population scale. The identification of autumn eels based on the otolith margin characteristics should be confirmed. It also indicates that the survival rate is very low for glass eels settling in the estuary in springtime.

REFERENCES


