Responses of basophilic cells of the digestive gland of mussels to petroleum hydrocarbon exposure

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ABSTRACT: A stereological study was conducted on the digestive gland of mussels Mytilus galloprovincialis Lmk., exposed to the water accommodated fraction of 2 crude oils and a commercial lubricant oil (3 different concentrations of each). Mussels were sampled after 21, 35, 77 and 91 d exposures, and the volume density (VD) of basophilic cells was determined in paraffin sections by a point-counting method. Exposure to petroleum hydrocarbons resulted in a significant dose-dependent increase in basophilic cell volume density. This was due to the presence of higher numbers of basophilic cells per digestive tubule, while the size of basophilic cells remained apparently unchanged. Results are interpreted in terms of an enhanced epithelial cell regeneration that might be linked to fundamental digestive processes. Indeed, a significant positive linear correlation exists between basophilic cell VD and the percentage of disintegrating tubules.

INTRODUCTION

The digestive diverticula of bivalves consist of numerous blind-ending tubules which communicate with the stomach by way of partially ciliated main ducts and non-ciliated secondary ducts (Owen 1970). The light microscope reveals that digestive and basophilic cells comprise the epithelium which lines the digestive tubules. Digestive cells are responsible for the intracellular digestion of food material while basophilic cells are thought to be involved in the synthesis and secretion of enzymes for extracellular digestion (Owen 1972). Early observations by Owen (1970) on the ultrastructure of the digestive tubules of Cardium edule revealed 2 different types of basophilic cells: pyramidal cells rich in granular endoplasmic reticulum and flagellated columnar cells. The later are considered to be immature and may serve to replace either the pyramidal basophilic cells, or the digestive cells, or both (Owen 1970). A similar cell type has been described in Mytilus edulis, held in the laboratory for long periods by Thompson et al. (1974), who suggest that the flagellate cell may differentiate into a mature digestive cell under suitable conditions.

Changes in molluscan digestive tubule structure due to pollutants have been studied quantitatively by Lowe et al. (1981), Tripp et al. (1984), Marigómez et al. (1986), Axiak et al. (1988), Recio et al. (1988) and Vega et al. (1989). Special attention was paid to the effects of pollutants on the structure and function of digestive cells since their lysosomal system is involved in hydrocarbon uptake (Moore 1980) and shows a high degree of responsiveness to alterations in the environment (Lowe et al. 1981, Harrison & Berger 1982, Moore et al. 1987, Cajaraville et al. 1989b, Marigómez et al. 1989). However, recent studies clearly suggest that basophilic cells are also subjected to important alterations under stress conditions. Petroleum hydrocarbon exposure induces the accumulation of large quantities of lipids in basophilic cells of mussels together with the fragmentation and vacuolation of cisterns of the rough endoplasmic reticulum and Golgi bodies (Carles et al. 1986, Lowe 1988). Further, Rasmussen et al. (1983) note that 'chronic' treatment of mussels with a N-nitroso compound causes varying degrees of degeneration in cells comprising the digestive epithelium, and that some digestive tubules are only composed of basophilic cells exhibiting dilatation and condensation of the endoplasmic reticulum. Widdows et al. (1984) found that exposure to petroleum hydrocarbons causes an 'apparent' increase in the numbers of basophilic cells in the prosobranch gastropod Littorina littorea while they do not report similar changes in Mytilus edulis maintained under the same conditions. An increased occurrence of basophilic cells has also been reported by us to occur in L. littorea under experimentally induced stress conditions.
conditions (Cajaraville et al. 1990, Marigomez et al. 1990). However, none of the studies mentioned above have applied quantitative techniques to measure changes in basophilic cell size or numbers; thus, the question of whether altered environmental conditions cause an increased presence of basophilic cells in molluscan digestive gland remains unanswered.

The purpose of the present study is to quantify the volume density of basophilic cells of the digestive gland of Mytilus galloprovincialis by using a stereological method and to describe the effects of the exposure to the Water Accommodated Fraction (WAF) of 3 different petroleum-derived hydrocarbons (PHC) on the basophilic cell volume.

MATERIAL AND METHODS

Experimental procedure. Mussels, collected from Meñakoz, Biscay, Spain (43° 24' N, 2° 93' W) in March 1988, were transferred to the laboratory. Fifty individuals (2.5 to 3.5 cm shell length) were distributed in 25 polystyrene-covered tanks in a thermostated semi-continuous water-flow system with activated charcoal and glass-wool filtered natural seawater (Zierbena, Biscay). Mussels were maintained unfed at 15 to 16 °C for 10 d to facilitate aclimatization to laboratory conditions. Subsequently, individuals were exposed in replicated series for 3 mo to different dilutions of the WAF of 2 crude oils (Maya and Ural types, supplied by Petronor SA, Spain), and a commercial lubricant oil (Repsol HD, made by Repsol SA, Spain).

The WAF of the oils was prepared using methods of Boylan & Tripp (1971) and Anderson et al. (1974). Oil (800 ml) was placed over 9 to 10 1 seawater in a 15 l Mariotte glass bottle with a rubber stopper and stirred with a magnetic stirrer for 12 h at room temperature, 18 to 20 °C. Stirring speed was adjusted to prevent the vortex of the mixture from extending more than 25% of the mixture’s height from the surface. WAF from each oil was produced twice every 2 d, exposure-time, type of toxicant) on the VD of basophilic cells. Correlation and regression analyses were performed to determine the effects of the factors studied (exposure-concentration, exposure-time, type of toxicant) on the VD of basophilic cells. Correlation and regression analyses were performed to establish significant relationships between basophilic cell VD and the above mentioned factors. In addition, correlation analyses were performed to establish significant correlations between changes in basophilic cell VD and changes in the percentages of the different digestive tubule types. For this purpose, digestive tubules were analysed using a subjective tubule grading method (Cajaraville et al. 1987a) where tubules were classified, according to

\[
VD = \frac{X_1 + X_2 + \ldots + X_n}{m \times n}
\]

where \(X = \) number of segment edges falling on basophilic cells; \(m = \) total number of segment edges falling on digestive tissue; \(n = \) number of counts (10 for each mussel).

Statistics. The statistical packet SPSS/PC+ (SPSS Inc., Microsoft Co.) was used in an AT personal computer (Atlas 286). Confidence intervals were calculated using Student's t-test and significant differences between means were established at the \(p < 0.05\) level (Sokal & Rohlf 1979). Significance of results was tested using a 3-way analysis of variance (ANOVA) to detect the effects of the factors studied (exposure-concentration, exposure-time, type of toxicant) on the VD of basophilic cells. Correlation and regression analyses were also carried out to establish significant relationships between basophilic cell VD and the above mentioned factors. In addition, correlation analyses were performed to establish significant correlations between changes in basophilic cell VD and changes in the percentages of the different digestive tubule types. For this purpose, digestive tubules were analysed using a subjective tubule grading method (Cajaraville et al. 1987a) where tubules were classified, according to
Langton (1975) into holding, absorbing, disintegrating and reconstituting. A 5th tubule type was considered necrotic (Cajaraville et al. 1989a).

RESULTS

Basophilic cells of *Mytilus galloprovincialis* can easily be distinguished from digestive cells in H & E stained sections under the light microscope (Fig. 1). Volume density (VD) of basophilic cells did not change in control mussels through the experimental period, except on Sampling day 35 where the value dropped markedly (Fig. 2).

The exposure to WAF of the 3 different PHC causes an increase in the VD of basophilic cells (Fig. 2). The response is observed in mussels exposed to ID and HD of the 3 PHC types tested and, with some exceptions, a dose-dependent effect is recorded. Values of mussels exposed to LD of Maya and Ural crude-oil types are not different from control values, while individuals exposed to LD of lubricant oil respond after 49 d exposure. VD values corresponding to HD of refined oil are missing on Sampling days 77 and 91 since mussels subjected to this treatment died after 49 d exposure. We recorded 100% mortality also in one of the ID series after 77 d exposure. Non-significant mortalities were noted during the whole experimental period in

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**Fig. 1. Mytilus galloprovincialis.** (A) Paraffin section of the digestive gland of a control individual sampled after 91 d maintenance in the laboratory; H & E 200×. (B) Higher magnification of the digestive tubules of the same individual; H & E 400×. (C) Paraffin section of the digestive gland of a mussel exposed for 91 d to the ID of the WAF of lubricant oil. Note that some tubules are comprised mostly of basophilic cells (arrows); H & E 200×. (D) Higher magnification of digestive tubules of the same individual; H & E 400×. T: digestive tubule, D: digestive cell, B: basophilic cell
mussels exposed to any dose of the 2 crude oils tested. In mussels treated with Maya, no mortalities occurred at LD. 1 mussel died after 21 d exposure to ID, and 4 died at HD (Days 50, 69, 70 and 74). In mussels treated with Ural, 1 individual died at LD (Day 79), 5 at ID (Days 37, 49, 71, 72 and 79) and 0 at HD.

Considering changes in VD of basophilic cells over the experimental period in the different treatments revealed that the lower values occurred on the 35th sampling day (as in the controls). Differences between control and treated mussels were always evident for each sampling period.

The most pronounced increase in basophilic cell VD was recorded in mussels treated with HD of the Ural-type crude oil after 21, 35 and 91 d exposure (19.13, 54.67 and 26.45% increase over control values from the respective sampling periods). An increase in the size of individual basophilic cells was not apparent in treated individuals. The increased VD recorded in PHC-exposed mussels is thus believed to be due to increased numbers of basophilic cells per digestive tubule (Fig. 1C, D). This was not observed in all digestive tubules of treated individuals, the response being limited to a number of tubules distributed through the whole digestive gland. This response pattern is the cause of the high variability found in stereological analyses between different fields within the same individual.

The 3-way analysis of variance (Table 1) indicates that there is a significant (p < 0.0001) effect of exposure time and exposure concentration on this parameter. In contrast, neither the type of PHC used nor the second- and third-order interactions between exposure time, exposure concentration and type of toxicant had a significant effect on the VD of basophilic cells. When interaction terms were suppressed to analyse the pure effect of each factor, the levels of significance remained unchanged. Correlation and regression analyses performed for each type of toxicant separately (Table 2) demonstrate that the VD of basophilic cells (in logits) increases linearly with increasing concentrations (in logits) of the toxicants. It must be stressed that regression coefficients do not vary significantly between types of toxicants as could be expected from ANOVA results. The variable exposure time is only significant in Ural-type crude oil. The regression equation for the total experimental population is similar to that of the Ural-type crude oil (Table 2). The dependent variable, Table 1 Three-way ANOVA. T: exposure-time, D: exposure-concentration, W: type of toxicant. Significance level = 95%
Table 2. *Mytilus galloprovincialis*. Regression equations of basophilic cell VD (log) against exposure-concentration (D), exposure-time (T) and concentration x time interaction (with and without logarithmic transformation) for each PHC tested and for the total experimental population. Only independent variables (VAR) with significant regression coefficients (p<0.05 according to Student's t-test) shown. K: Y-intercept; b: regression coefficient; SE(b): standard error of b; p(b): significance of b based on Student's t-test; r: multiple correlation coefficient; p(r): significance of r based on Fisher's F-test; df: degrees of freedom.

<table>
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<tr>
<th>Ural</th>
<th>VAR</th>
<th>b</th>
<th>SE(b)</th>
<th>p(b)</th>
<th>r</th>
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<td>0.0543</td>
<td>0.0006</td>
<td>0.3979</td>
<td>0.0001</td>
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<tr>
<td>T</td>
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<td>0.0114</td>
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<td>K</td>
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<td>0.2723</td>
<td>0.0059</td>
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<tr>
<td>K</td>
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<td>0.0224</td>
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<table>
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<th>SE(b)</th>
<th>p(b)</th>
<th>r</th>
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<tr>
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<td>0.3258</td>
<td>&lt;0.00001</td>
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<tr>
<td>T</td>
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VD of basophilic cells (×100), was log-transformed since regression equations developed with untransformed values showed lower significance of regression coefficients. Correlation analyses between basophilic cell VD and percentages of each tubule type are shown in Table 3. Both linear and logarithmic correlation coefficients were calculated. Correlation coefficients with the best significance for each pair basophilic cell VD/tubule type percentage have been indicated (Table 3). In mussels exposed to the WAF of the Ural-type crude oil, basophilic cell VD increases with increasing percentages of disintegrating tubules and decreasing percentages of absorptive tubules (in logits), while the VD of...
basophilic cells (in logits) is negatively correlated with the percentage of holding tubules. For mussels exposed to the Maya-type crude oil, VD (in logits) covaries negatively with both the percentage of holding tubules and the percentage of necrotic tubules. For the lubricant oil, basophilic cell VD is positively correlated with the percentage of disintegrating tubules. Taking into account that the ANOVA indicates no significant differences in basophilic cell VD between different PHC, the correlation analysis was performed for the whole experimental population. It then became apparent that basophilic cell VD is correlated positively with the percentage of disintegrating tubules and negatively with the percentage of necrotic tubules and the percentage of absorbing tubules (in logits). Basophilic cell VD (in logits) is also negatively correlated with the percentage of holding tubules. In no case was a significant correlation recorded between basophilic cell VD and percentage of reconstituting tubules.

DISCUSSION

The volume density of basophilic cells was quantified by stereology in the digestive gland of *Mytilus galloprovincialis*. Both control and treated mussels showed very low values of basophilic cell VD on the 35th sampling day (early May). An explanation for this phenomenon could be obtained from a parallel study carried out under identical experimental conditions (Cajaraville et al. 1989a). In this study a slight increase in both digestive tubule epithelial thickness and digestive tubule size was recorded in mussels (controls and experiments) at the 35th sampling day. These changes could account for a reduced probability of finding basophilic cells in a given microscopic field. Besides, these results seem to be related to the reproductive state of mussels since the lowest values of the gonad index (Seed 1969) occur, especially in females, also in this sampling period (own unpubl. results). Durfort (1989) finds that the progression of the different developmental stages of oocytes is associated with changes in digestive gland Type B cells, involved in calcium storage and mobilisation, in the polyplacophoran mollusc *Acanthochitona fascicularis*. However, seasonal variations in digestive and basophilic cell numbers of digestive tubules and in digestive tubule size have, to our knowledge, never been reported; thus this idea is rather speculative.

Significant alterations were recorded in basophilic cell VD as a result of petroleum hydrocarbon exposure. The 3 different PHCs tested induce a significant increase in basophilic cell VD due to the presence of increased numbers of basophilic cells per digestive tubule. Both hydrocarbon exposure concentration and exposure time exert a significant effect on the VD of basophilic cells. There was no significant interaction between concentration and time, indicating that increased values of the parameter were maintained throughout the experiment without any long-term recovery or deterioration. Besides, there was no significant effect of the type of toxicant used (2 different crude oils and 1 lubricant oil) which clearly indicates the result is independent of the PHC tested. In fact, the regression analyses carried out separately for each toxicant show that basophilic cell VD increases with increasing exposure concentrations after double logit transformation. This further implies an attenuated slope in the response at lower and higher exposure concentrations, a linear dose dependence being achieved only when mussels were exposed to intermediate concentrations. Nevertheless, although the 3 PHCs evoked a similar response pattern in basophilic cells it is evident that the lubricant oil is much more toxic to mussels since it is the only toxicant causing significant mortality. It is also the only toxicant inducing an increase in basophilic cell VD at the low exposure dose. In other studies, WAF of refined oils was reported to be more toxic than WAF of crude oils (Anderson et al. 1974, Rice et al. 1976).

Widdows et al. (1984) found an apparent increase in the number of basophilic cells in *Littorina littorea* exposed to petroleum hydrocarbons, while a 2.5 d recovery was sufficient to cause a reduction in basophilic cell numbers. Since an increased incidence of basophilic cells has also been suggested in studies concerning 1-naphthol and cadmium toxicities in *L. littorea* (Cajaraville et al. 1990, Marigomez et al. 1990), it could be hypothesized that the response is general among molluscs and independent of the stressor involved. These characteristics, together with the above-reported significant dose-dependent effect, could allow a potential use of VD of basophilic cells as a general stress indicator in marine pollution-monitoring programs. Furthermore, quite distinct stress sources such as crowding of the larval stages of *Schistosoma japonicum* in the intertubular spaces of the digestive gland of the gastropod *Oncomelania hupensis quadraasi* resulted in a marked increase of pyramidal calcium cells (possibly equivalent to bivalve basophilic cells) together with vacuolation of digestive cells (Querubin & Enriquez 1989). This again seems to indicate that basophilic cells of molluscan digestive tissue respond to a variety of harmful conditions by increasing their numbers and that this response is usually associated with degenerative changes in digestive cells (Thompson et al. 1974, Rasmussen et al. 1983, Querubin & Enriquez 1989, Cajaraville et al. 1990, Marigomez et al. 1990). Whether increased numbers of basophilic cells are simply due to concomitant reduction in digestive
cell numbers or whether basophilic cell proliferation is a necessary step for digestive cell regeneration remains to be investigated. Studies by Mix & Sparks (1971) and Thompson et al. (1974) suggest that replacement of digestive cells occurs by division of (a type of) basophilic cell. Thus, proliferation of basophilic cells could be related to an increased loss of digestive cells which would require an increased cell turnover and regeneration in damaged digestive tubules.

In context with this hypothesis, an interesting question is whether different tubular types (see Langton 1975 for a description of phasic types of digestive tubules) bear different numbers of basophilic cells. Preliminary results (present paper) indicate that high values of basophilic cell VD occur when the percentage of disintegrating tubules in a given digestive gland is also high (and subsequently, percentages of holding, absorbing and necrotic tubules are low). In no case was the basophilic cell VD significantly correlated with the percentage of reconstituting tubules. Thus, changes in basophilic cell numbers could be associated with phasic activities of the digestive gland and consequently, to both digestion process and turnover of digestive cells. Widdows et al. (1984) consider the PHC-induced increase in basophilic cell numbers to be an adaptive response in order to meet the increased demands on the digestive secretory system resulting from the presence of oil. In conditions in which digestive cells are challenged, and thus the intracellular digestion process is affected, an increase in basophilic cell numbers might be necessary to augment enzyme secretion for extracellular digestion (Marigomez et al. 1990). This again suggests that alterations in basophilic cell incidence could be related to fundamental digestive cell processes.

More information is needed on both the role of basophilic cells in bivalve digestive gland physiology and the reactions of these cells to the presence of environmental PHC. At the cellular level, fragmentation and vacuolation of the rough endoplasmic reticulum, dilation of the Golgi apparatus (Carles et al. 1986) and accumulation of lipid droplets (Carles et al. 1986, Lowe 1988) occur. Whether these alterations in basophilic cells are a consequence of hydrocarbon metabolism and detoxification is not known. Triebkorn (1989) suggests a potential involvement of calcium cells of terrestrial slugs in the metabolism of carbamates and metaldehyde since an intensified activity of NADPH-neotetrazolium reductase was observed in basophilic cells of Littorina littorea (Moore et al. 1982) where blood cells display the strongest reactivity. Thus, involvement of basophilic cells in detoxication of products deriving from PHC exposure seems unlikely; their increased numbers may not reflect enhancement of detoxication processes.

In conclusion, exposure to sublethal WAF concentrations of petroleum-derived hydrocarbons results in a significant, dose-dependent increase in VD of basophilic cells of the digestive gland of mussels. These changes could be closely linked to the loss of digestive cells, and to a subsequent wave of basophilic cell proliferation leading to a replacement of the cells lost. Studies are currently being conducted to enable a more rapid quantification of this parameter by automated image analysis.

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