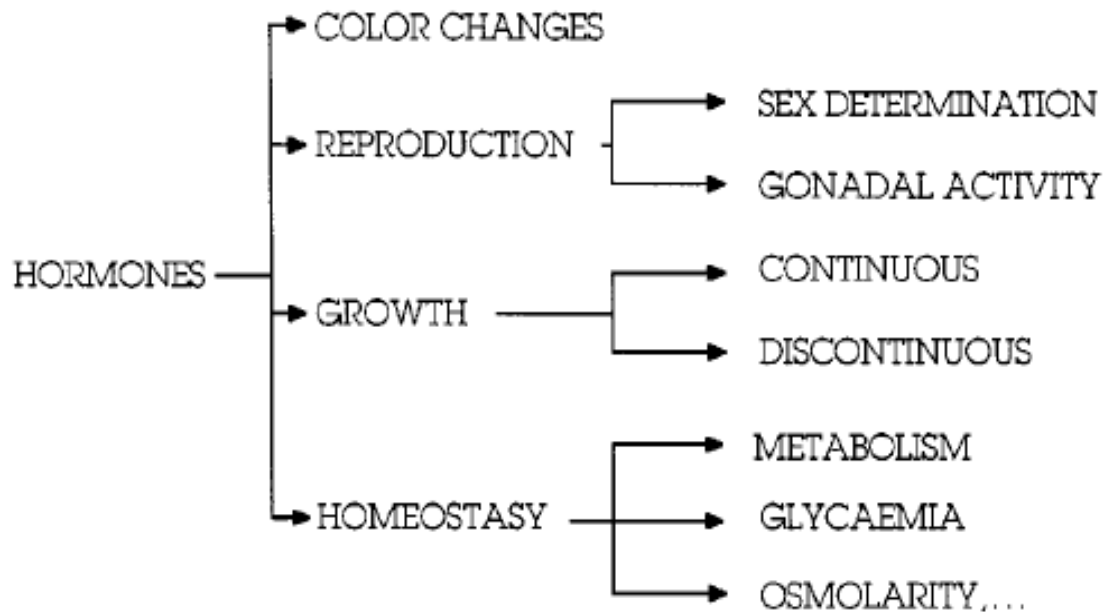


# Endokrinní disrupce - bezobratlí

## Hormonální regulace u bezobratlých:

- oblast intenzivního výzkumu - praktické aspekty ochrany zemědělských plodin; endokrinní disrupce jako faktor ovlivňující stabilitu ekosystémů;
- přehledné publikace jsou poměrně zastaralé - především primární zdroje;
- otázka definice - endokrinní orgány (Mollusca) vs. endokrinní buňky (Annelida)

# Hormonální regulace u bezobratlých:



**neuropeptidy vs. klasické hormony**

## Hormonální regulace u bezobratlých:

TABLE I

Examples of Reported Hormones in Different Invertebrate Taxa<sup>a</sup>

| Taxon        | Reported hormones (example, <i>controlled process</i> )   |
|--------------|---|
| Coelenterata | <b>Neuropeptides</b> (glycine-leucine tryptophan amides = GLWamides, <i>metamorphosis</i> ); thyroids (thyroxine, <i>strobilation</i> ); retinoids (9-cis-retinoic acid, <i>strobilation</i> )  |
| Nematoda     | <b>Ecdysteroids</b> (reported but <i>functional role questionable</i> ); <b>terpenoids</b> (juvenile hormone (JH) like hormones, <i>growth</i> ); neuropeptides (FMRFamide, <i>function unknown</i> )   |
| Mollusca     | Ecdysteroids (reported but <i>functional role questionable</i> ); steroids (17 $\beta$ -estradiol, testosterone, progesterone, <i>sexual differentiation, reproduction in prosobranchs</i> ); terpenoids (JH reported but <i>functional role questionable</i> ); neuropeptides (APGWamide, dorsal body hormone (DBH), <i>sexual differentiation, gonad maturation, spawning</i> ; egg-laying hormone (ELH), <i>spawning</i> ; FMRFamide, <i>neuromodulation</i> ; molluscan insulin-like peptides (MIPs), <i>growth, development, energy metabolism</i> ) |
| Annelida     | Ecdysteroids (ecdysone, <i>functional role unknown</i> ); neuropeptides (FMRFamide, <i>neuromodulation</i> )  |

## Hormony u členovců:

### Crustacea

Ecdysteroids (ecdysone, *molting, vitellogenesis*); steroids (17 $\beta$ -estradiol, testosterone, progesterone, *functional role under debate*); terpenoids (methyl farnesoate (MF), *metamorphosis, reproduction*); neuropeptides (androgenic hormone, *sexual differentiation, vitellogenesis inhibition*; crustacean hyperglycemic hormone family (CHH), *energy metabolism*; molt-inhibiting hormone (MIH), *ecdysteroid production*; vitellogenesis-inhibiting hormone (VIH), *vitellogenesis*)

### Insecta

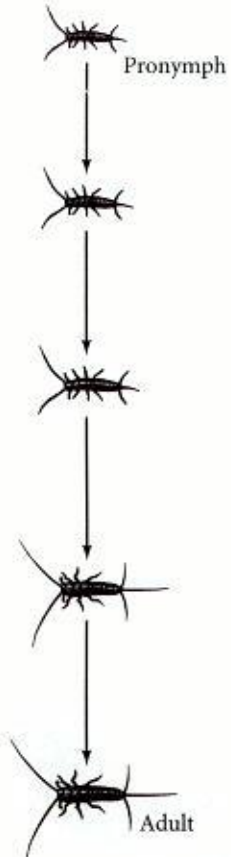
Ecdysteroids (ecdysone, *molting, egg maturation*); terpenoids (JH, *metamorphosis, reproduction*); neuropeptides (adipokinetic hormone (AKH), *energy metabolism*; allatostatin and allatotropin, *JH production*; bombyxin, *ecdysteroid production, energy metabolism*; bursicon, *cuticle tanning*; diapause hormone, *embryonic diapause*; diuretic hormone (DH), *water homeostasis*; ecdysis-triggering hormone (ETH) and eclosion hormone (EH), *ecdysis behavior*; FMRFamides, *neuromodulation*; prothoraciotrophic hormone (PTTH), *ecdysteroid production*)

## Hormony u ostnokožců a pláštěnců:

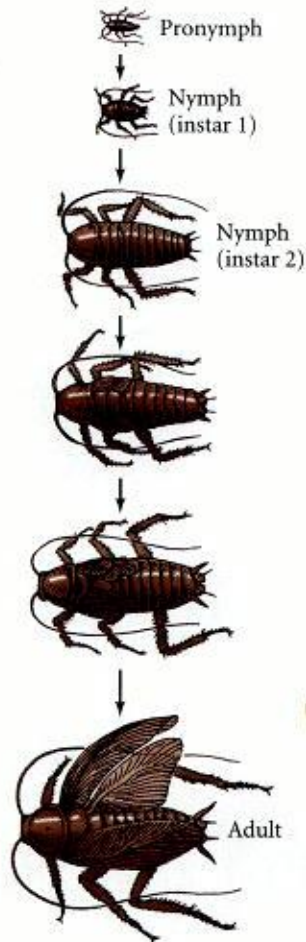
- Echinodermata **Steroids** progesterone, testosterone,  $17\beta$ -estradiol, estrone, *vitellogenesis*, *oogenesis*, *spermatogenesis*, *spawning*);  
neuropeptides (gonad-stimulating substance = GSS, *spawning*;  
maturation-promoting factor = MPF, *fertilisation*)
- Tunicata Steroids (testosterone,  $17\beta$ -estradiol, *oogenesis*, *spermatogenesis*,  
*spawning*); neuropeptides (gonadotropin releasing hormone  
analogue, *gonad development*); **thyroids** (thyroxine, *probably  
tanning process during tunic formation*)

## Metamorfóza hmyzu:

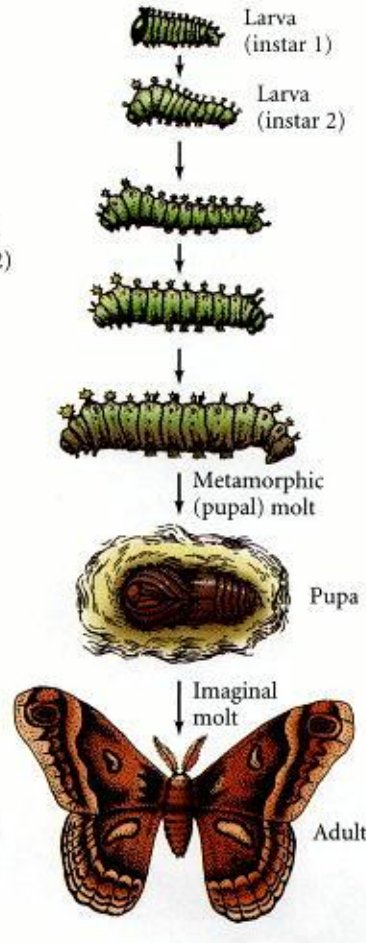
(A) AMETABOLOUS DEVELOPMENT



(B) HEMIMETABOLOUS DEVELOPMENT



(C) HOLOMETABOLOUS DEVELOPMENT



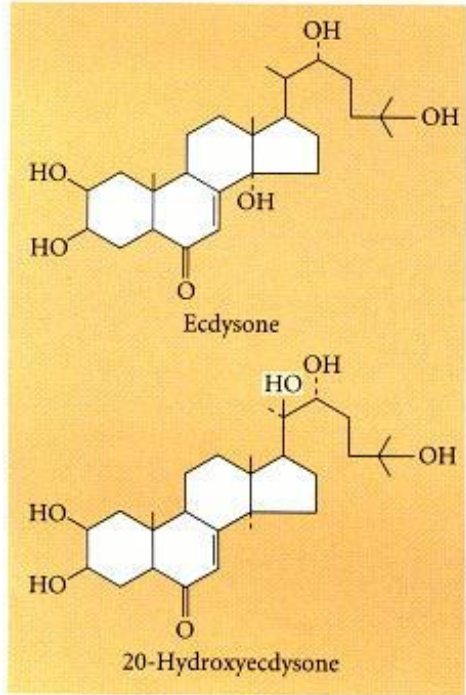
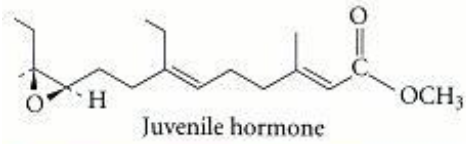
Modes of insect development. Molts are represented as arrows.

(A) Ametabolous (direct) development in a silverfish. After a brief pronymph stage, the insect looks like a small adult.

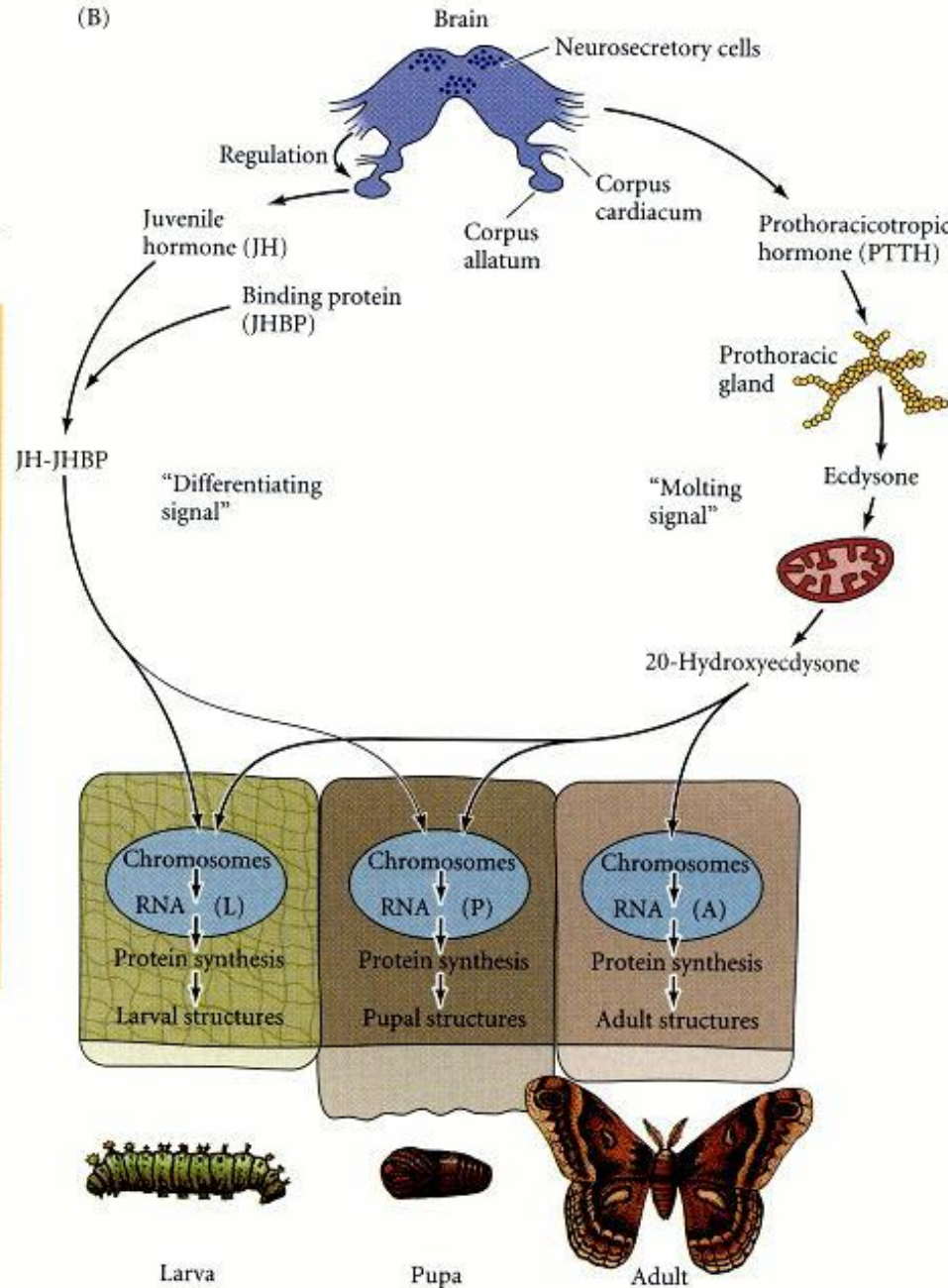
(B) Hemimetabolous (gradual) metamorphosis in a cockroach. After a very brief pronymph phase, the insect becomes a nymph. After each molt, the next nymphal instar looks more like an adult, gradually growing wings and genital organs.

(C) Holometabolous (complete) metamorphosis in a moth. After hatching as a larva, the insect undergoes successive larval molts until a metamorphic molt causes it to enter the pupal stage. Then an imaginal molt turns it into an adult.

(A)

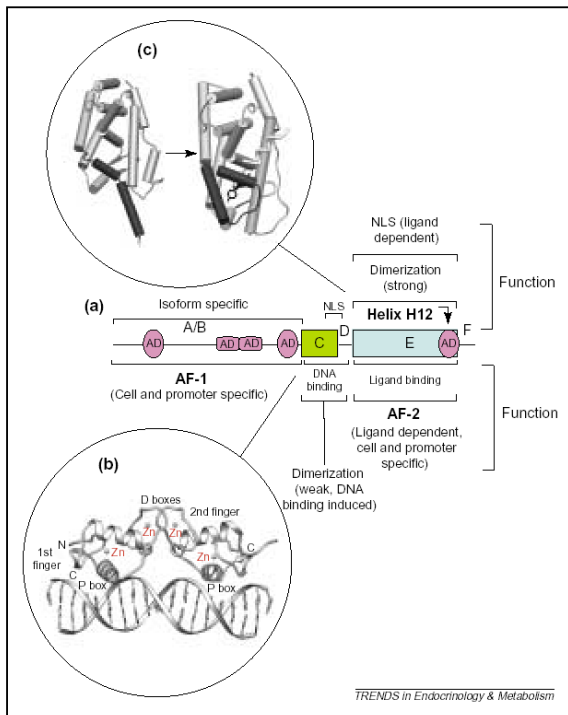


(B)

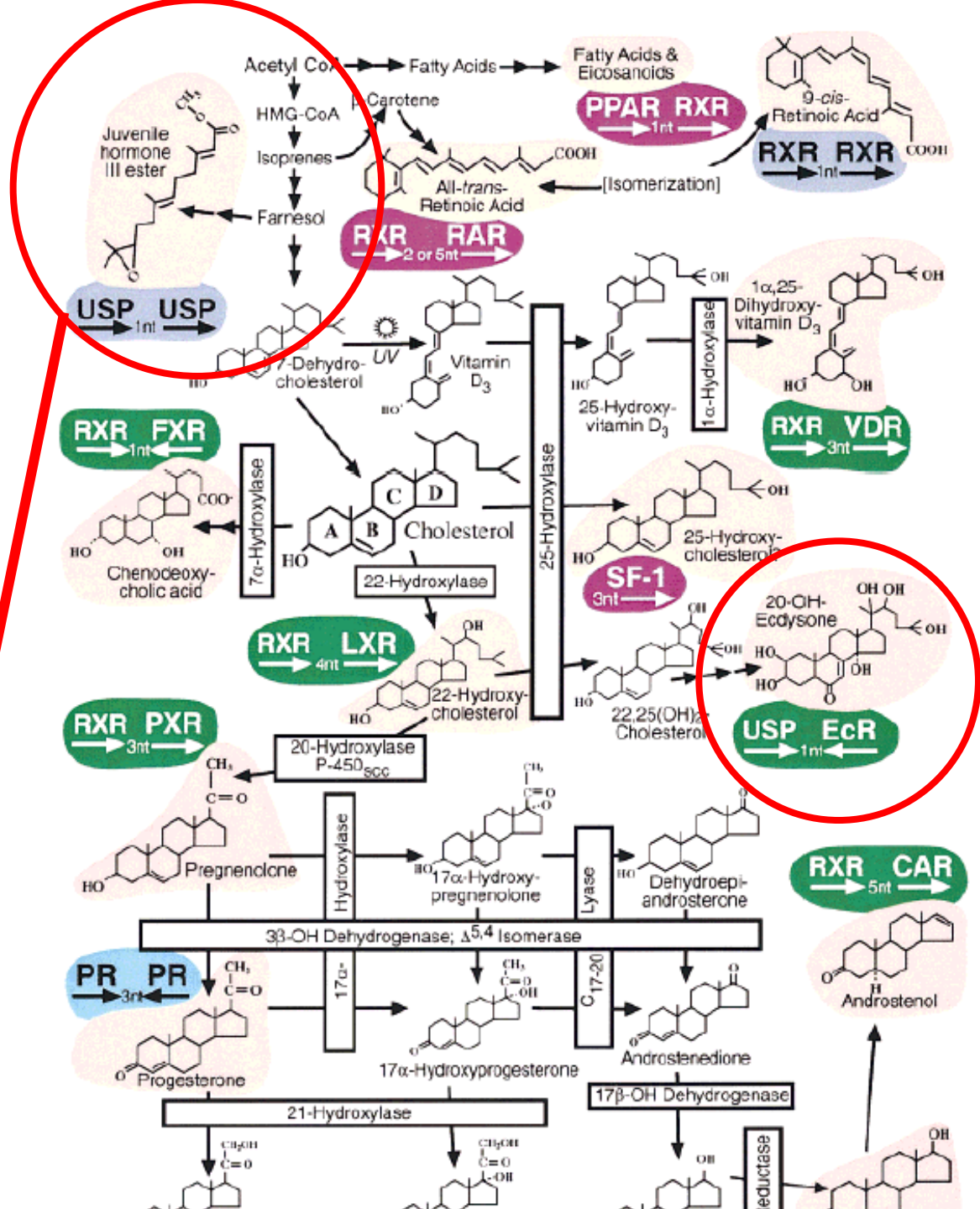


Regulation of insect metamorphosis. (A) Structures of juvenile hormone, ecdysone, and the active molting hormone 20-hydroxyecdysone. (B) General pathway of insect metamorphosis. Ecdysone and juvenile hormone together cause molts to keep the status quo and form another larval instar. When there is a lower concentration of juvenile hormone, the ecdysone-induced molt produces a pupa. When ecdysone acts in the absence of juvenile hormone, the imaginal discs differentiate, and the molt gives rise to the adult

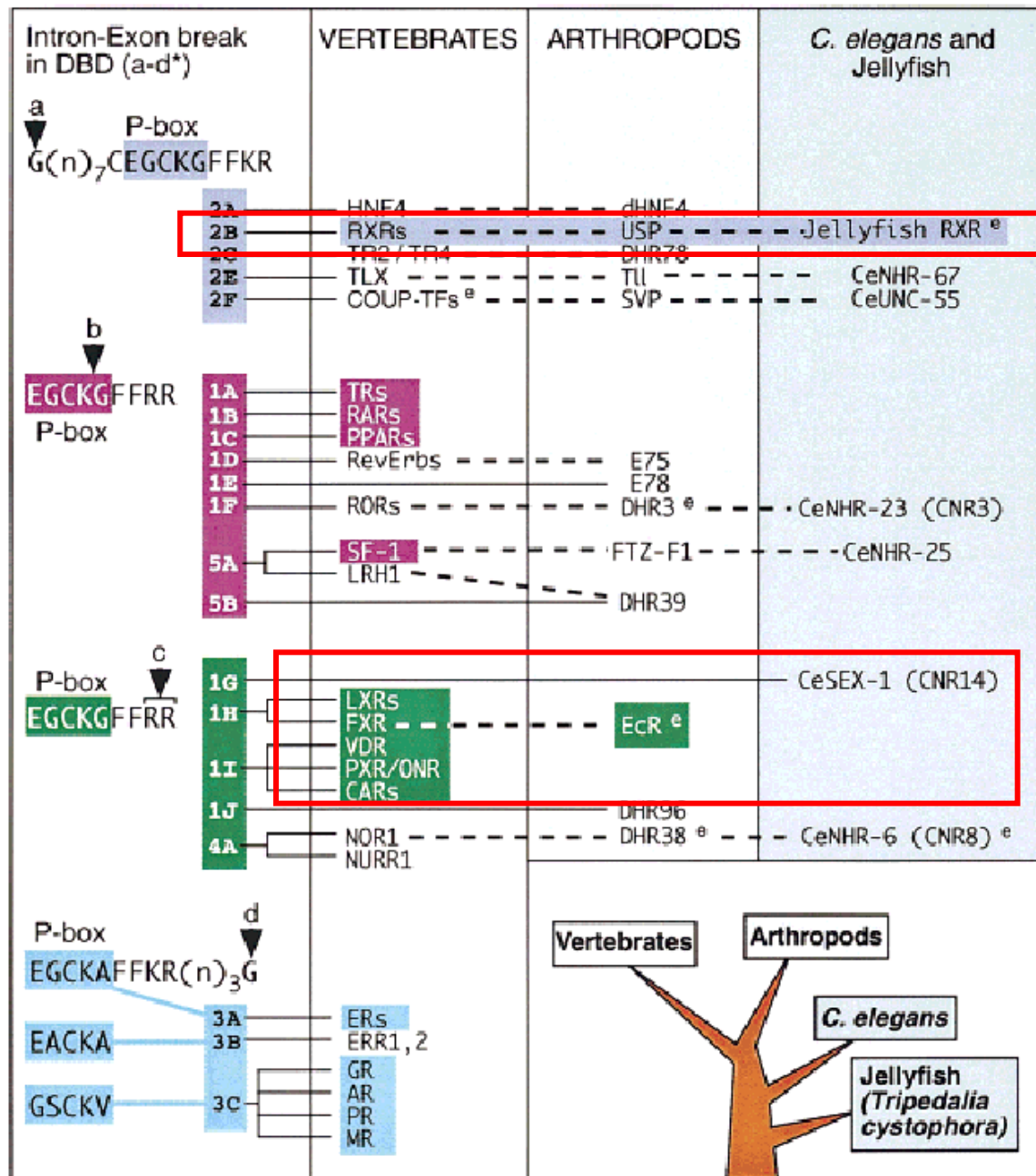




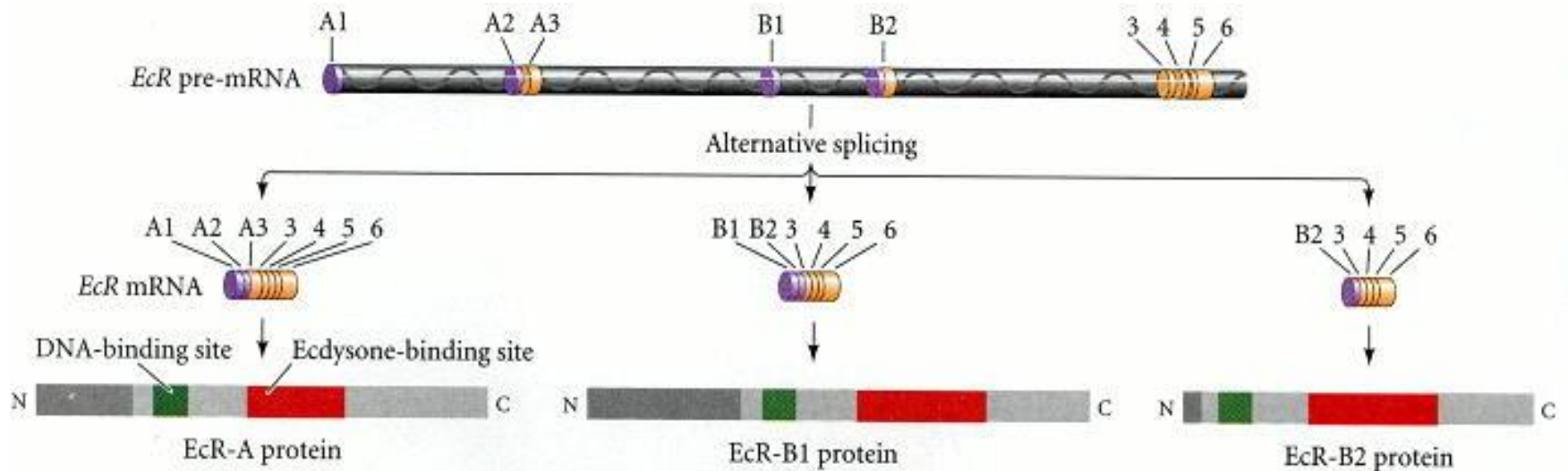
**Fig. 1.** (a) Schematic of the structural and functional organization of NRs. The evolutionary conserved regions C (DBD) and E (LBD) are indicated as boxes and a black line represents the divergent regions A/B, D and F. Two transcription AFs have been described in several NRs, a constitutively active (if taken out of the context of the receptor) AF-1 in region A/B and a ligand-inducible AF-2 in region E. Within these AFs, AFs have been defined. (b) Estrogen receptor DBD complex on a cognate DNA response element. (c) Agonist-induced changes of the LBD, allowing binding of coactivators (the bound coactivator-binding peptide is shown). Figures 1b,c are three-dimensional views derived from the corresponding crystal structures. Abbreviations: See Glossary.



**Another candidate for the JH receptor role is the Methoprene-tolerant (Met) Per-Arnt-Sim (PAS) domain protein, whose loss confers tolerance to JH and its mimic methoprene in the fruit fly *Drosophila melanogaster*. (Konopová et al., 2007.**



## Ec receptor:

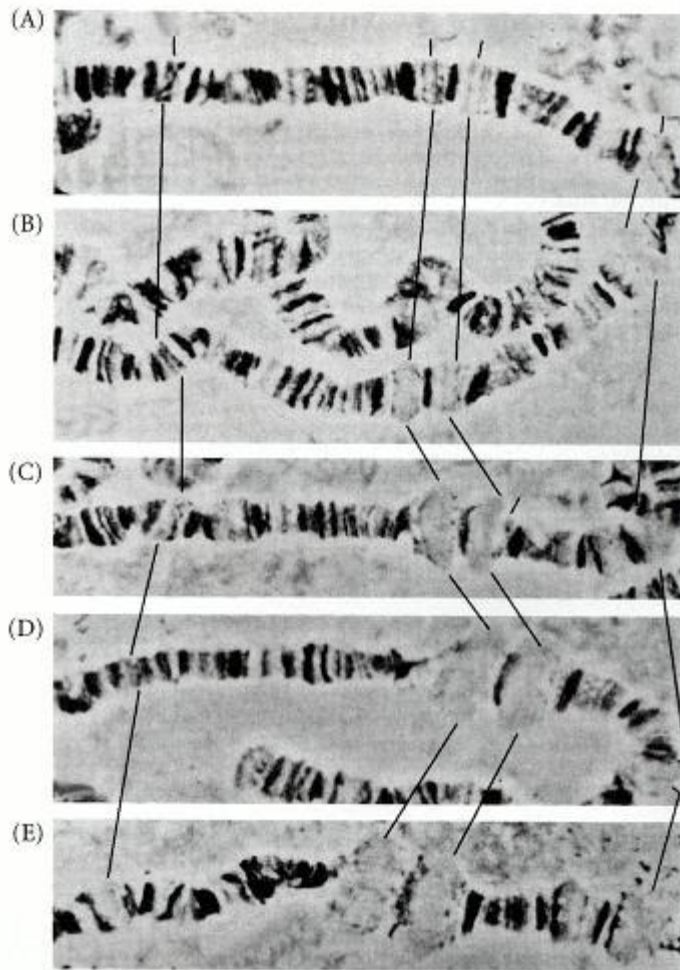


Formation of the ecdysone receptors. Alternative mRNA splicing of the ecdysone receptor (*EcR*) transcript creates three types of *EcR* mRNAs. These generate proteins having the same DNA-binding site (blue) and hydroxyecdysone-binding site (red), but with very different amino termini.

Three isoforms of EcR have been identified in insects, each with a different, stage-specific role in regulation of molting and development. This allows for one steroid hormone to induce a variety of different tissue responses. In general, EcR A is predominant when cells are undergoing a maturation response (from juvenile to adult) and is predominant in imaginal discs, whereas EcR B1 predominates in juvenile cells during proliferation or regression. Little is known about the function of the EcR B2 isoform.

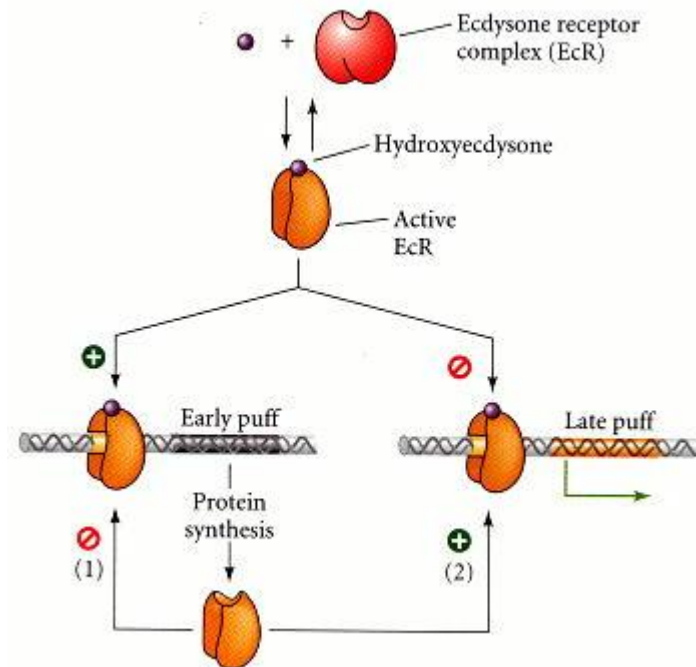
DNA and hormone binding are similar in the three isoforms of EcR. Little is known about the crustacean EcR isoforms and how they change during the molt cycle. However, the EcR that has been cloned from the crab, *Uca pugilator* (U31817, GenBank), shares 85-87% homology with that of *Drosophila* (M74078, GenBank). The differences are primarily in the region of the molecule involved with dimerization. Similar sequence similarities are found between the heterodimeric partner, USP.

There are several ecdysteroids which bind EcR, including 20-hydroxyecdysone, turkesterone, makisterone A, ponasterone A, and muristerone A. Some arthropods may use specific ecdysteroids as their principal molting hormone, but often several ecdysteroids are found within one group. The primary molting hormone for a range of organisms, including some insects and crustacea, is 20-OH ecdysone (20 HE). Among other examples, makisterone A is an important hormone for some crustacea and hemipteran insects.



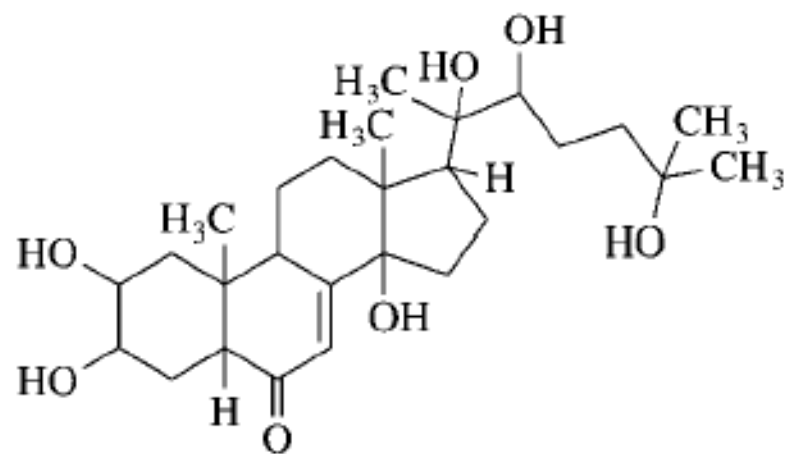
Hydroxyecdysone-induced puffs in cultured salivary gland cells of *D. melanogaster*. (A) Uninduced control. (B-E) Hydroxyecdysone-stimulated chromosomes at (B) 25 minutes, (C) 1 hour, (D) 2 hours, and (E) 4 hours.

The Ashburner model of hydroxyecdysone regulation of transcription. Hydroxyecdysone binds to its receptor, and this compound binds to an early puff gene and a late puff gene. The early puff gene is activated, and its protein product (1) represses the transcription of its own gene and (2) activates the late puff gene, perhaps by displacing the ecdysone receptor. (After [Richards 1992.](#))

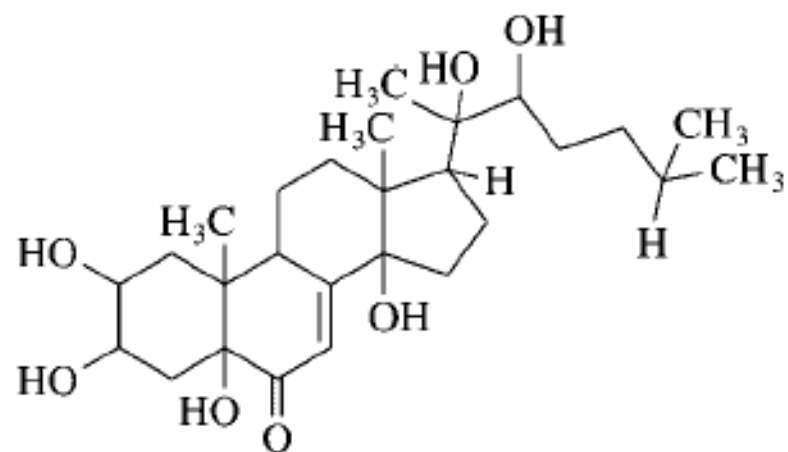


## Struktury Ecdysonů:

20-OH Ecdysone  
(20 HE)

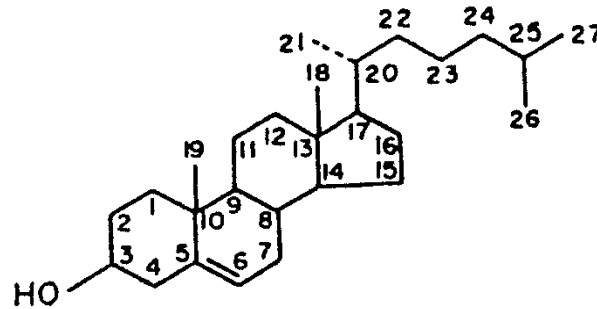


Muristerone A  
(MurA)



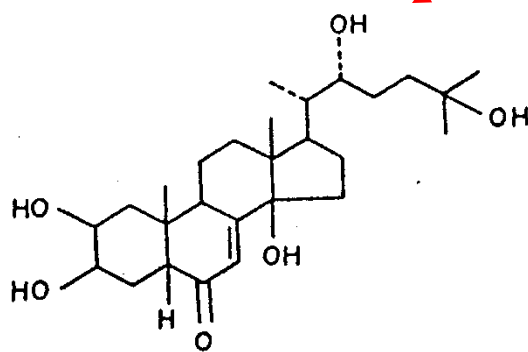
# Synthesis of molting hormones

Cholesterol (from diet- a vitamin for insects)



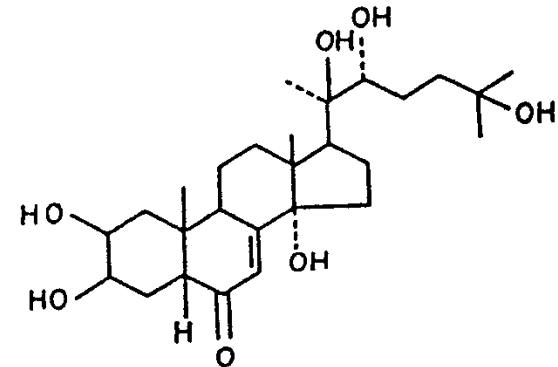
Prothoracic gland

Metabolites (excretion)  
Conjugates (storage)



Ecdysone

mono-oxygenase  
(fat body, epidermis)

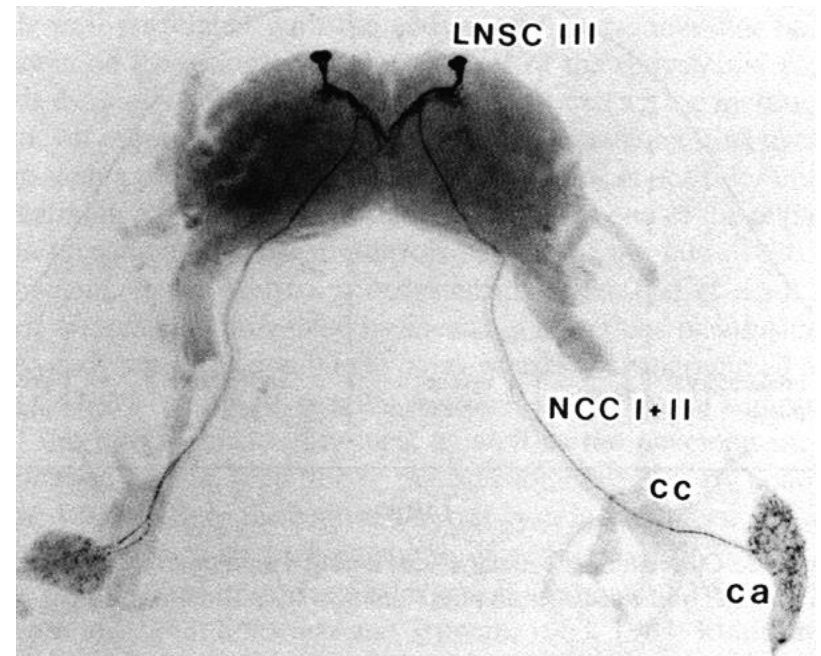


20-Hydroxyecdysone

## Prothoracicotropic hormone (PTTH)

- Protein of 30-kDa active as a homodimer linked by a disulfide bond
- Produced by two pairs of lateral neurosecretory cells
- Released from corpora allata (moths) or corpora cardiaca (most insects)

PTTH cells of *Manduca* visualized with anti-PTTH antibody →





# The role of PTTH

Ecdysone synthesis and secretion are initiated by prothoracicotropic hormone (PTTH), a hormone produced by two pairs of neurosecretory cells in the brain. PTTH was first isolated and characterized from the silkworm *Bombyx mori*. PTTH has conserved seven cysteine residues, several hydrophobic regions and an *N*-glycosylation site. Only the homodimeric form of *Bombyx* PTTH is biologically active. *Bombyx* PTTH is thought to be a member of the transforming growth factor- $\beta$  (TGF- $\beta$ ) family.

## ■ Initiates every molt

- stimulates prothoracic glands to synthesize and release ecdysone
- serves as “mission control” allowing molt if the conditions are right
  - factors affecting decision to molt are species-specific
    - stretch of the abdomen by a blood meal in *Rhodnius*
    - completion of the cocoon in some moths
    - escape from wet diet in flies

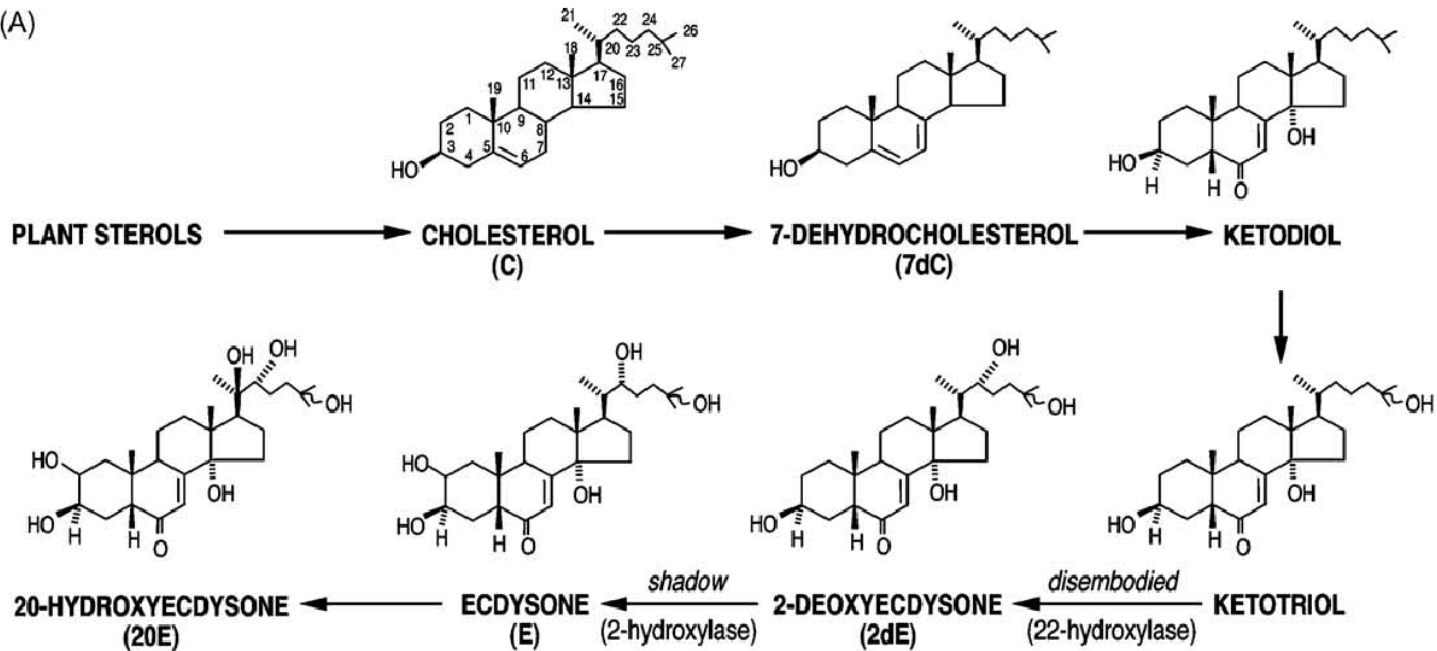
## Fytoekdosteroidy:

Phytoecdysteroids (PEs) are a family of about 200 plant steroids related in structure to the invertebrate steroid hormone 20-hydroxyecdysone. Typically, they are C27, C28 or C29 compounds possessing a 14 $\alpha$ -hydroxy-7-en-6-one chromophore and A/B-cis ring fusion (5 $\beta$ -H).

PEs are attracting renewed attention because of their specific effects on invertebrate development (potential in invertebrate pest control) and their varied benign pharmacological actions on mammals (biomedical applications and gene switches). In the past three decades, several thousand species of plants have been surveyed for the presence of PEs and the structures of over 200 PEs have been deduced. The most frequently encountered PE is 20E, the principal physiological inducer of moulting and metamorphosis in arthropods.

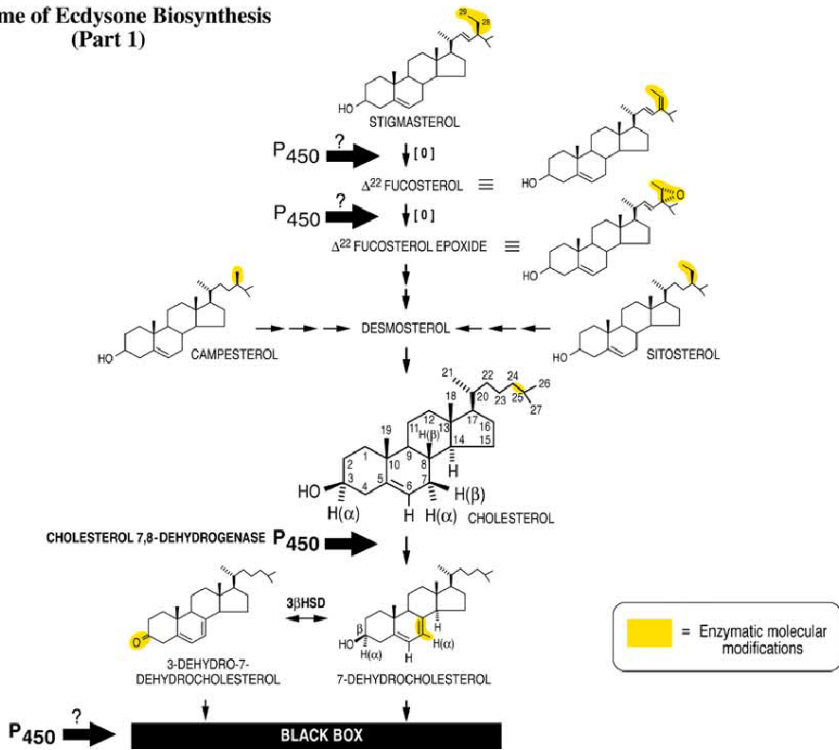
# Syntéza ekdysteroidů a úloha cytochromů P450:

(A)



# Syntéza ekdysteroidů a úloha cytochromů P450:

Scheme of Ecdysone Biosynthesis (Part 1)



Scheme of Ecdysone Biosynthesis (Part 2)

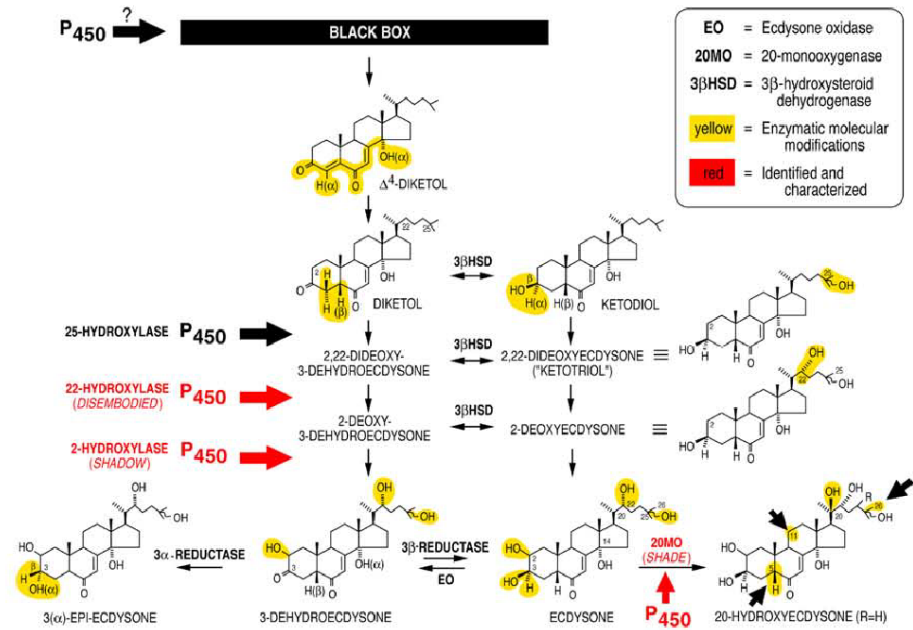
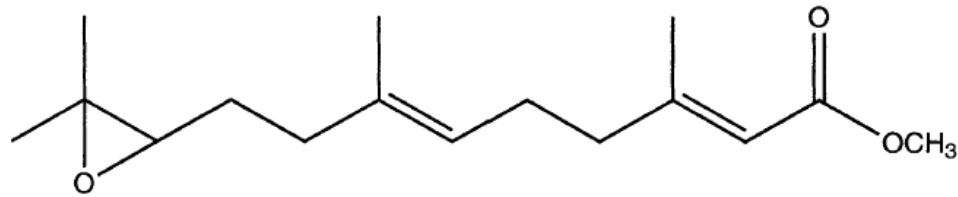
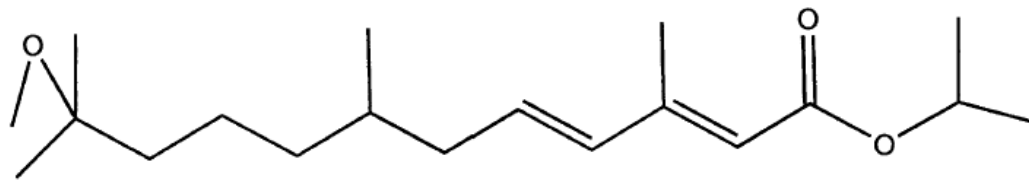


Fig. 5. (Parts 1 and 2). The biosynthesis of 20-hydroxyecdysone from plant sterols. Question marks denote possible involvement of P450 enzymes. Note specifically where the Halloween gene products act (red). 3-Dehydroecdysone is synthesized in the prothoracic glands of many insects (e.g. *Manduca sexta*) and converted to ecdysone in the hemolymph (left column of part 2). For *Drosophila*, ecdysone is synthesized in the prothoracic gland cells of the ring gland (right column of part 2).

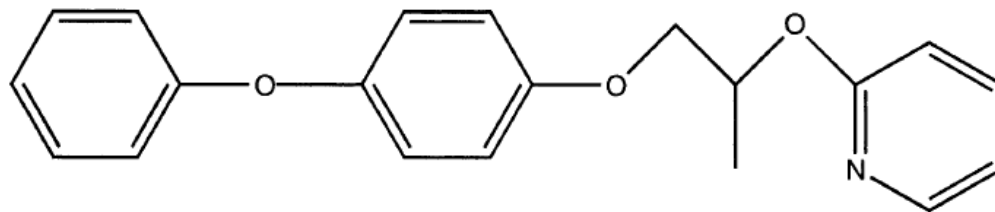
## Juvenilní hormon a jeho analogy:



JH III



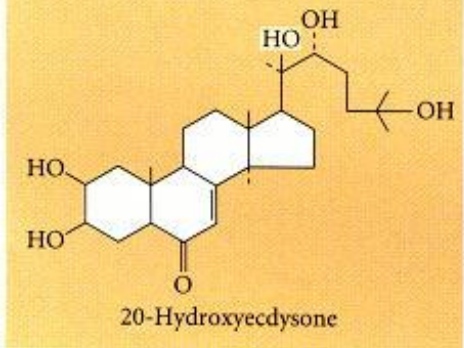
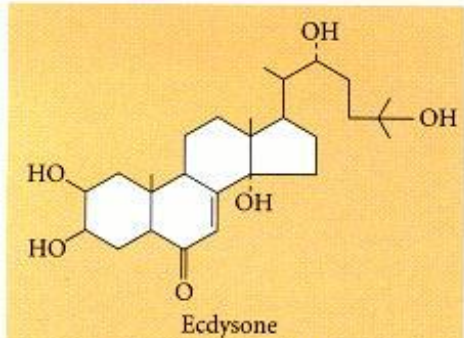
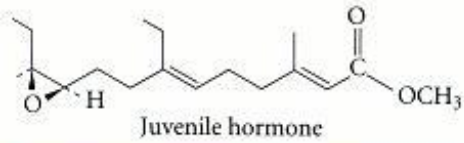
Methoprene



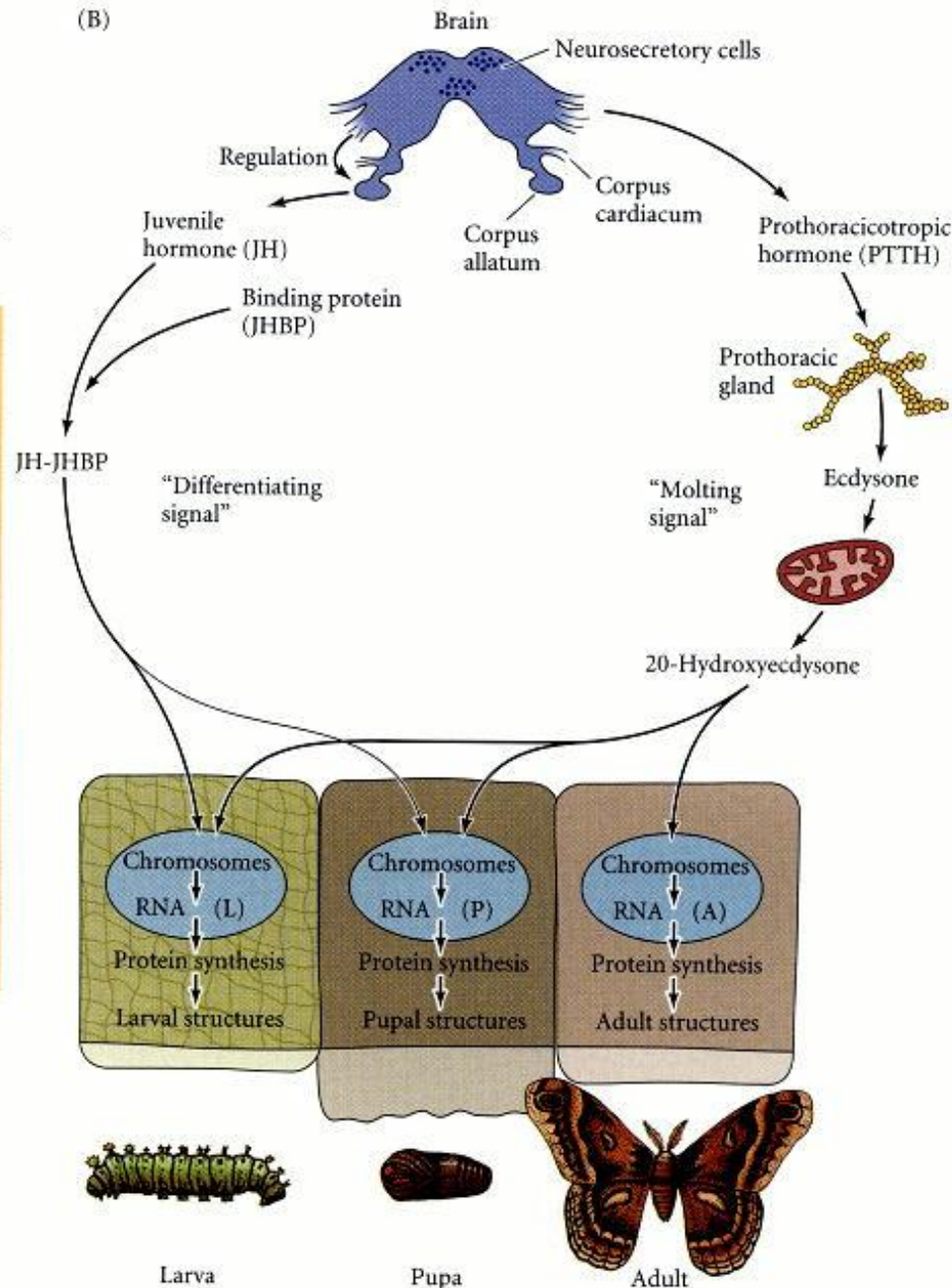
Pyriproxyfen

Fig. 1. Chemical structures of JH III and JHAs.

(A)



(B)



**Regulation of insect metamorphosis. (A) Structures of juvenile hormone, ecdysone, and the active molting hormone 20-hydroxyecdysone. (B) General pathway of insect metamorphosis. Ecdysone and juvenile hormone together cause molts to keep the status quo and form another larval instar. When there is a lower concentration of juvenile hormone, the ecdysone-induced molt produces a pupa. When ecdysone acts in the absence of juvenile hormone, the imaginal discs differentiate, and the molt gives rise to the adult**

Insect juvenile hormones are critical developmental hormones that have direct effects on both larval development and adult reproductive competence. Most insect orders appear to synthesize a single JH homolog, methyl (2*E*,6*E*)-10,11-epoxy-3,7,11-trimethyl-2,6-dodecadienoate (JH III) but Lepidoptera and at least some Diptera synthesize additional homologs.

Although not well documented, regulation of JH production is complicated and involves endogenous neural, neuroendocrine signals and, in some cases, male produced exogenous regulators transferred to the female during mating. Among virgin females, JH is required for vitellogenesis and, thus, females do not become reproductively competent until JH production is stimulated. To date, only two neuropeptides that regulate JH biosynthesis in adult Lepidoptera have been identified. These were identified e.g. from the tobacco hornworm moth (*Manduca sexta*) and are:

**allatotropin**

(Gly-Phe-Lys-Asn-Val-Glu-Met-Met-Thr-Ala-Arg-Gly-Phe-NH<sub>2</sub>)

**allatostatin**

(pGlu-Val-Arg-Phe-Arg-Gln-Cys-Tyr-Phe-Asn-Pro-Ile-Ser-Cys-Phe-COOH).

*Manduca sexta*





INSECT PHEROMONE

# Insect Pheromone Biochemistry and Molecular Biology

## The biosynthesis and detection of pheromones and plant volatiles

This book provides an up-to-date and in-depth coverage of how insects produce pheromones and how they then detect both pheromones and plant volatiles. Well over half the species on the planet are insects - more than 800,000 in all. Their perception of each other and their world is achieved through the production and reception of chemical odors that provide essential information for the location of prospective mates and food supplies. These chemical messages are unique for each species and thus represent a vast landscape in which we may explore the evolution of behavior and communication.

Many insects such as moths, beetles, aphids and grasshoppers are also pests of crops. Attracted by the specific smells released by these plants, they not only find food, but also each other, aided further by the odorous pheromones that they synthesize and release. Feeding and breeding are thus equally served by their extraordinary sense of smell. Understanding the underlying mechanisms of odor detection and pheromone biosynthesis offers us the means to disrupt their predations and populations without the use of harmful and poisonous pesticides. In the realm of disease transmission, insects such as mosquitoes, ticks and fleas feed on the blood of humans and other animals and in so doing transfer the dangerous pathogens which cause illnesses such as malaria, lime disease and plague. As with the plant feeders, these insects find their hosts by smell. Again, understanding the underlying mechanism of odor detection can help us to combat the process and contribute to improving human health. Continuing research into insect olfaction, founded on the study of insect pheromones, thus provides tremendous scope for mitigating the profound socio-economic impact of insects.

Gary Blomquist and Richard Vogt have been leaders in pheromone production and reception, respectively, for over 20 years. Blomquist was co-editor of the very successful 1987 book "Pheromone Biochemistry", and has organized and been part of numerous symposia on the biosynthesis and endocrine regulation of pheromone production. Vogt was involved in the pioneering work on the biochemistry and molecular biology of pheromone reception in insects and has remained a leader in this area. He has organized and participated in numerous symposia and written several reviews on the subject.



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Insect Pheromone Biochemistry  
and Molecular Biology

Blomquist • Vogt



# Insect Pheromone Biochemistry and Molecular Biology

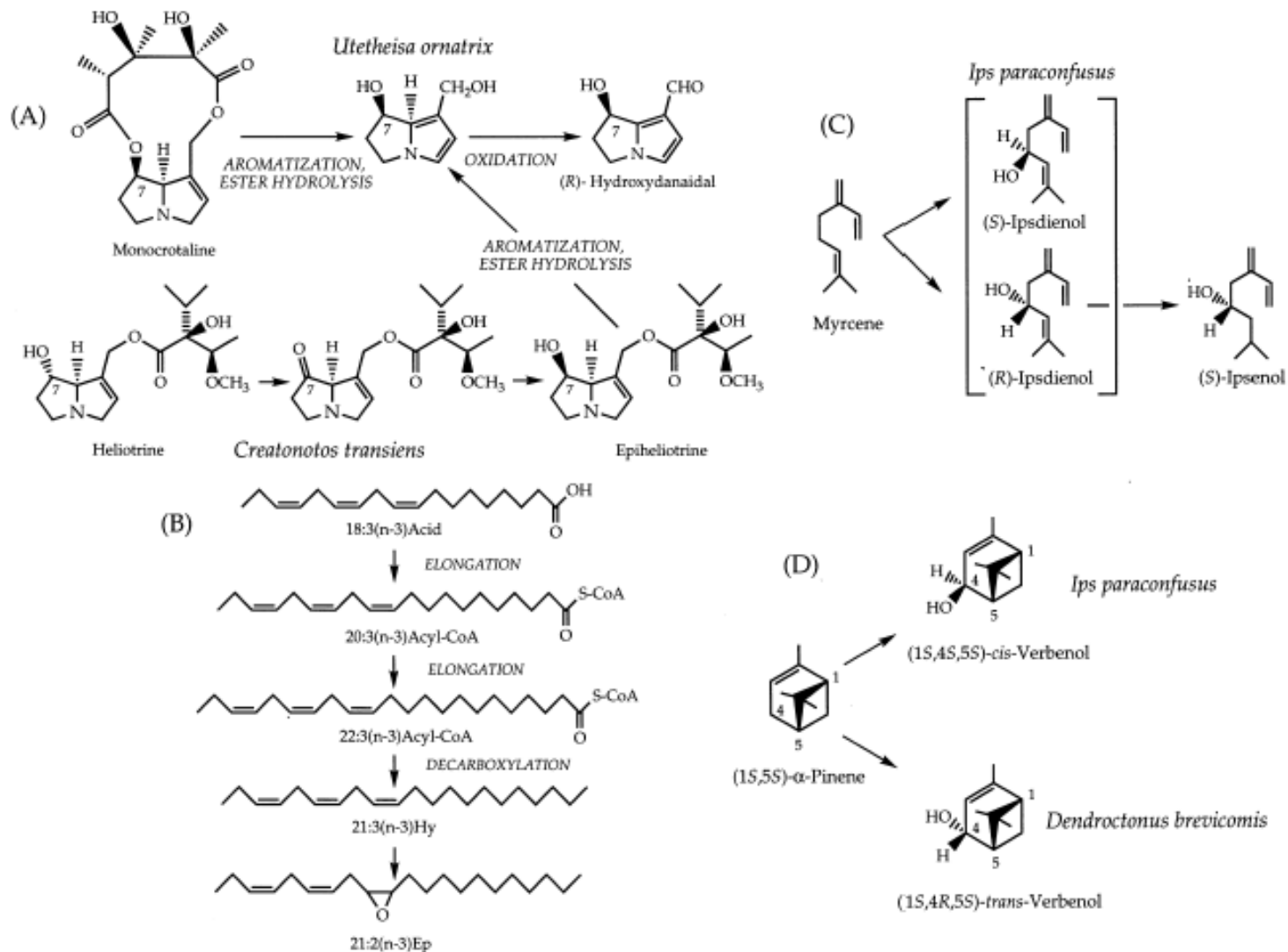
## The biosynthesis and detection of pheromones and plant volatiles



Gary J. Blomquist  
Richard G. Vogt



# Feromony jsou strukturně vysoce odlišné látky



# Syntéza feromonů může být přímo regulována hormony

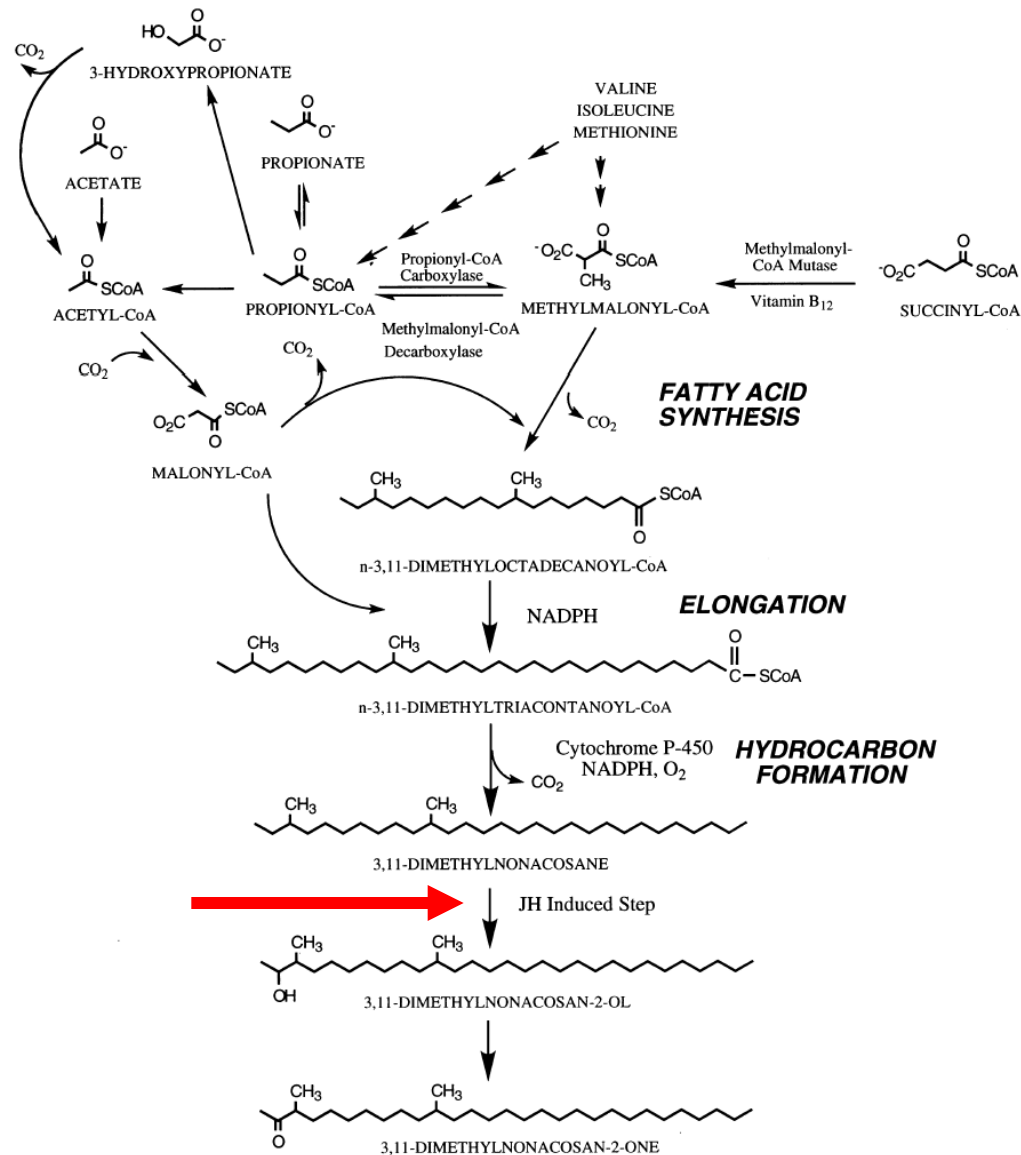
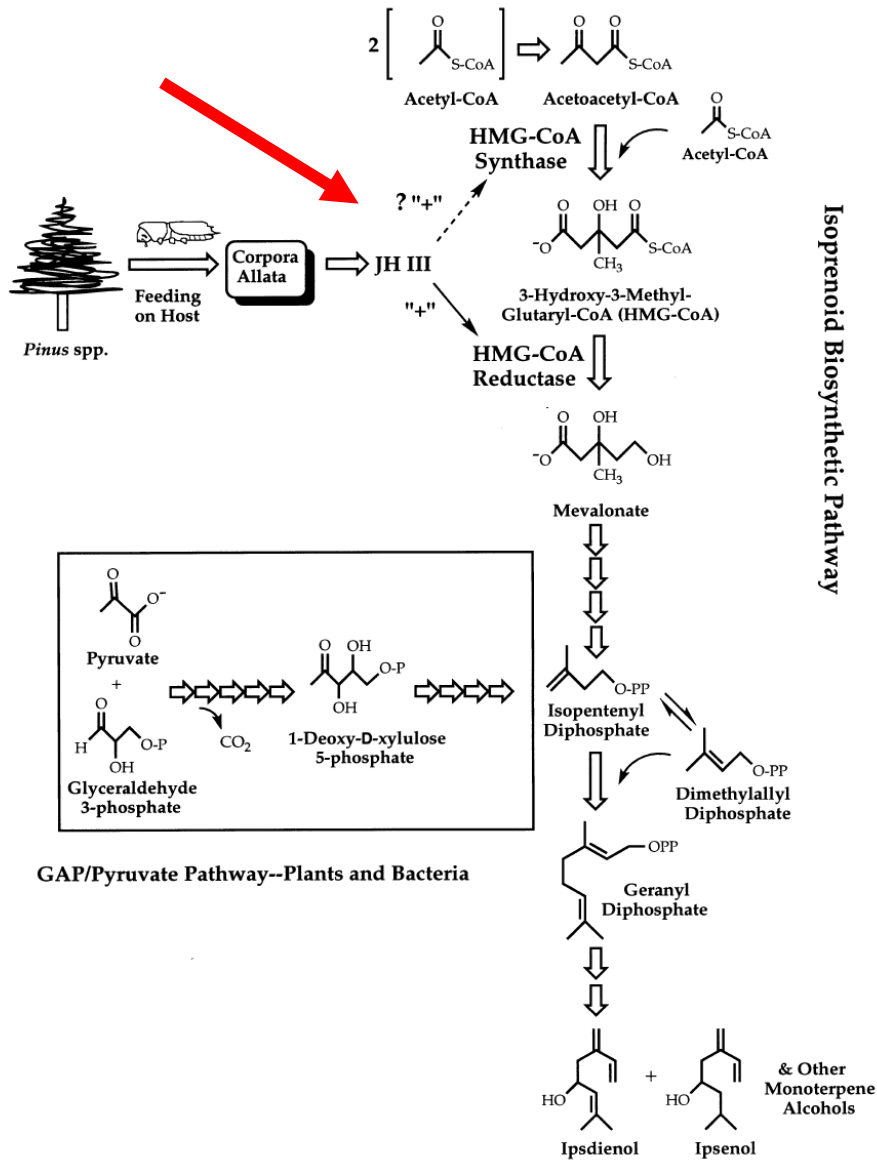


Fig. 2. Blattodean pheromone biosynthetic pathways utilize fatty acid biosynthesis from malonyl-CoA and methylmalonyl-CoA substrates followed by cytochrome P-450-mediated decarboxylation, hydroxylation, and oxidation. The hydroxylation step is regulated by JH III (adapted from Chase et al., 1992 for *Blattella germanica* sex pheromone components).



Coleopteran pheromone biosynthetic pathways as exemplified for *Ips* spp. [e.g. *Ips pini* (Say)] and acyclic monoterpene (ipsdienol) pheromone biosynthesis. The classical mevalonate-based isoprenoid pathway is regulated by juvenile hormone III (JH III) at enzymatically catalyzed steps prior to mevalonate. Feeding on host *Pinus* spp. phloem induces synthesis of JH III by the corpora allata.

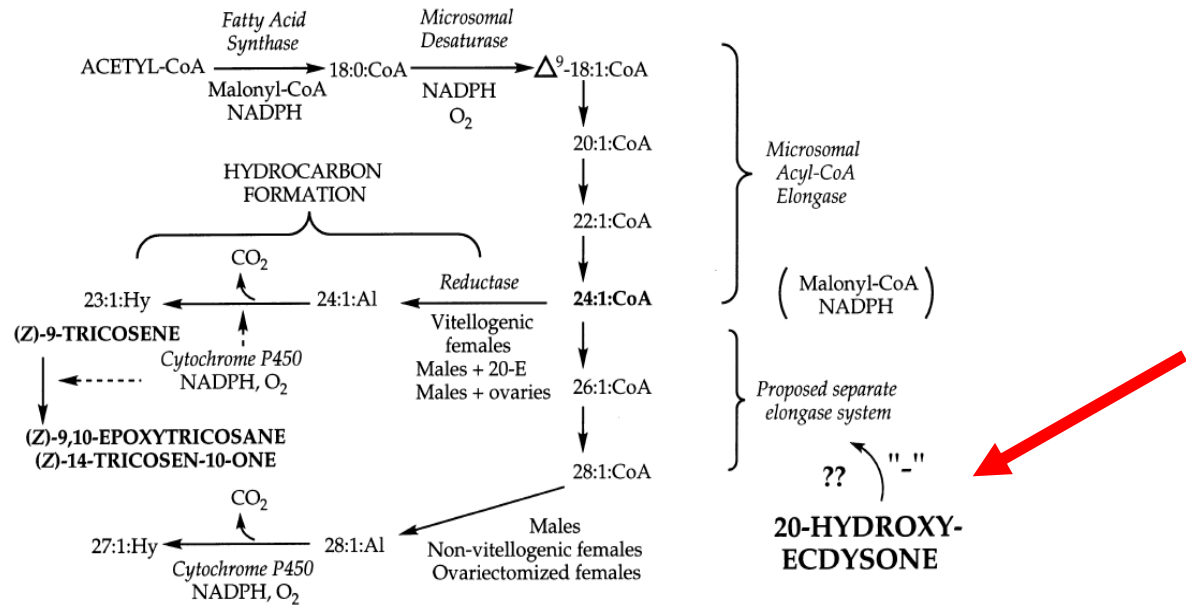


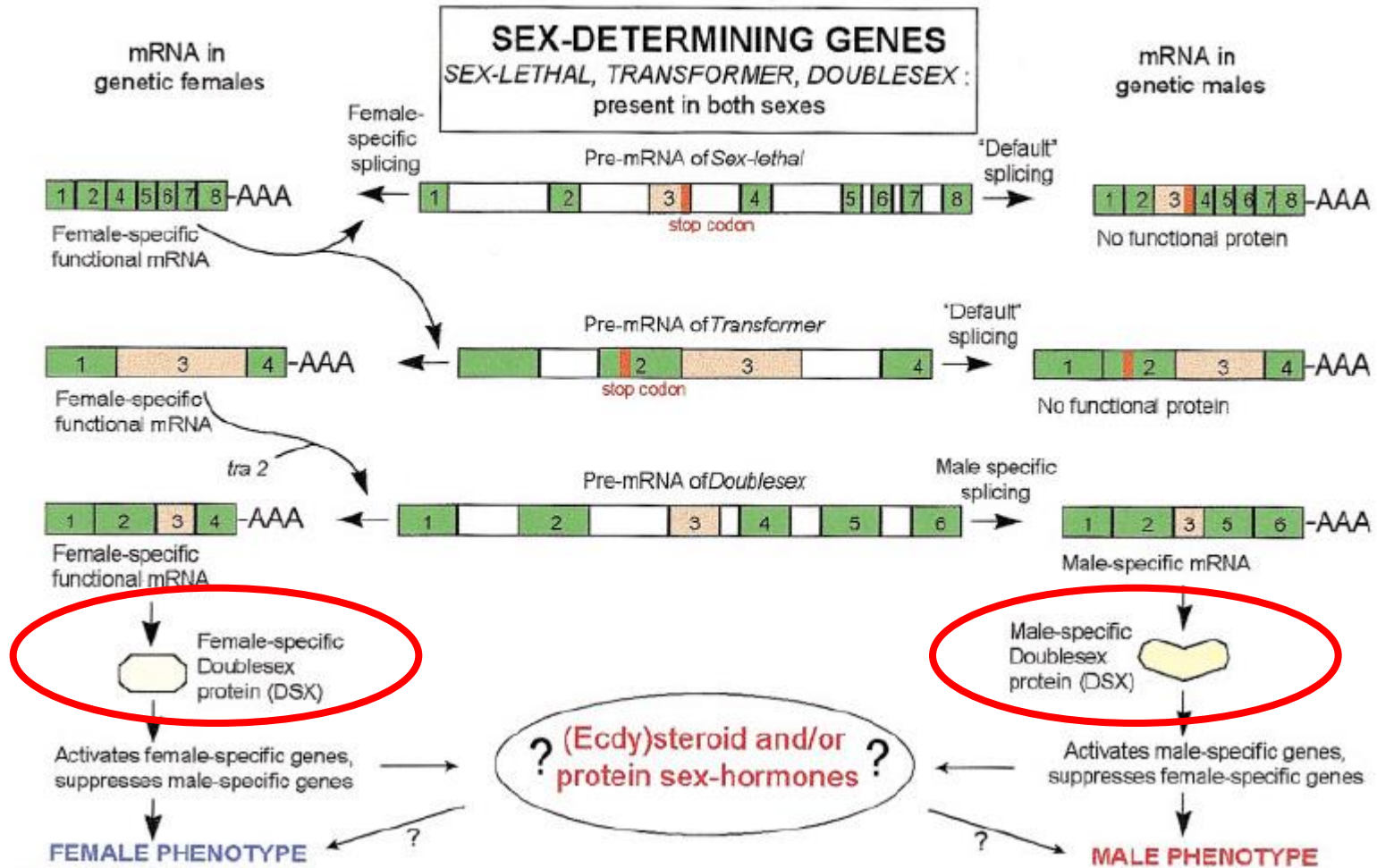
Fig. 4. Dipteran pheromone biosynthetic pathways utilize fatty acid synthesis, desaturation, elongation, and reductive decarboxylation. The proposed regulatory steps for 20-hydroxyecdysone are the secondary elongation system. Unsaturated hydrocarbons can be further modified to the epoxides (adapted from Blomquist et al., 1987a for the common house fly, *Musca domestica* L. sex pheromone components).

# Vyskytují se u hmyzu pohlavní hormony?

FEMALE GENOTYPE : AA + XX

DROSOPHILA

MALE GENOTYPE : AA + XY



## Endokrinní disrupce u bezobratlých:

The issue of endocrine disruption (ED) in invertebrates has generated remarkably little interest in the past compared to research with aquatic vertebrates in this area. However, with more than 95% of all known species in the animal kingdom, invertebrates constitute a very important part of the global biodiversity with key species for the structure and function of aquatic and terrestrial ecosystems. Despite the fact that ED in invertebrates has been investigated on a smaller scale than in vertebrates, invertebrates provide some of the best documented examples for deleterious effects in wildlife populations following an exposure to endocrine-active substances. The principal susceptibility of invertebrates to endocrine-active compounds is demonstrated with the case studies of tributyltin effects in mollusks and of insect growth regulators, the latter as purposely synthesized endocrine disrupters.

## Imposex:

The first adverse effects of TBT on mollusks were observed in *Crassostrea gigas* at the Bay of Arcachon, one of the centers of oyster aquaculture in Europe with ball-shaped shell deformations in adults, and a dramatically decline of annual spatfall. These effects led to a break-down of local oyster production in the bay with marked economic consequences. Laboratory and field analyses revealed that TBT from antifouling paints was the causative agent with trace concentrations as low as 10 to 20 ng TBT/L in ambient water being already effective. Another TBT effect in mollusks was initially described in a number of regions worldwide in the early 1970s without identifying the organotin compound as the causative agent at that time: A virilization of female prosobranchs, which has been termed as **imposex**. **Imposex is characterized by the formation of a penis and/or vas deferens on females of gonochoristic prosobranch species and is induced at lower concentrations than all other described TBT effects.** Furthermore, it is a specific response of organotin compounds under field conditions. Today, imposex is known to occur in more than 150 prosobranch species.



## Imposex:

Tributyltin (TBT)

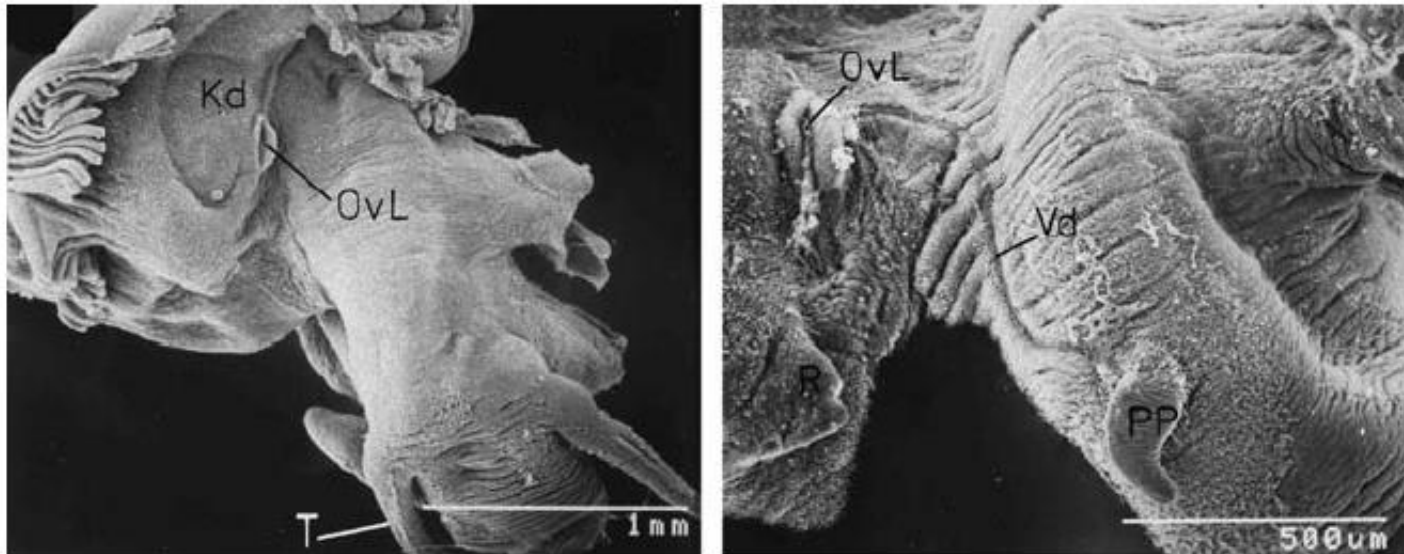
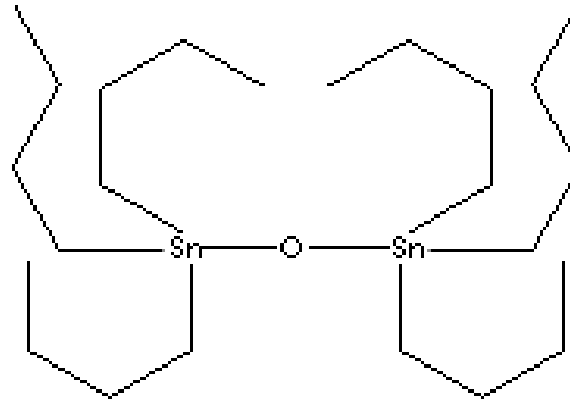


Fig. 1 *Hydrobia ulvae*. Scanning electron micrographs of females with their mantle cavity opened. Left: normal female without imposex; right: sterilized female in the final stage of imposex with blocked oviduct. Abbreviations: Kd, capsule gland; OvL, Ooporous opening of oviduct (open left; blocked right); PP, Penis; T, tentacle; Vd, vas deferens.

The periwinkle *Littorina littorea* develops a closely related virilization phenomenon as a response to TBT exposure, termed as **intersex**. Intersex females are either characterized by male features on female pallial organs, specifically by an inhibition of the ontogenetic closure of the pallial oviduct or female sex organs are supplanted by the corresponding male formations particularly by a prostate gland. Comparably to imposex, the intersex response is a gradual transformation of the female pallial tract, which can be described by an evolutive scheme with four stages.

Intersex development causes restrictions of the reproductive capability of females. In stage 1, a loss of sperm during copulation is possible and consequently the reproductive success is reduced. Females in stages 2-4 are definitively sterile because the capsular material is spilled into the mantle cavity (stage 2) or the glands responsible for the formation of egg capsules are missing (stages 3 and 4). Due to female sterility, periwinkle populations can be in decline but are not likely to become extinct because of the planktonic veliger larvae produced by the species, as long as aqueous TBT levels are not beyond mortality threshold concentrations for the larvae (Matthiessen *et al.* 1995).

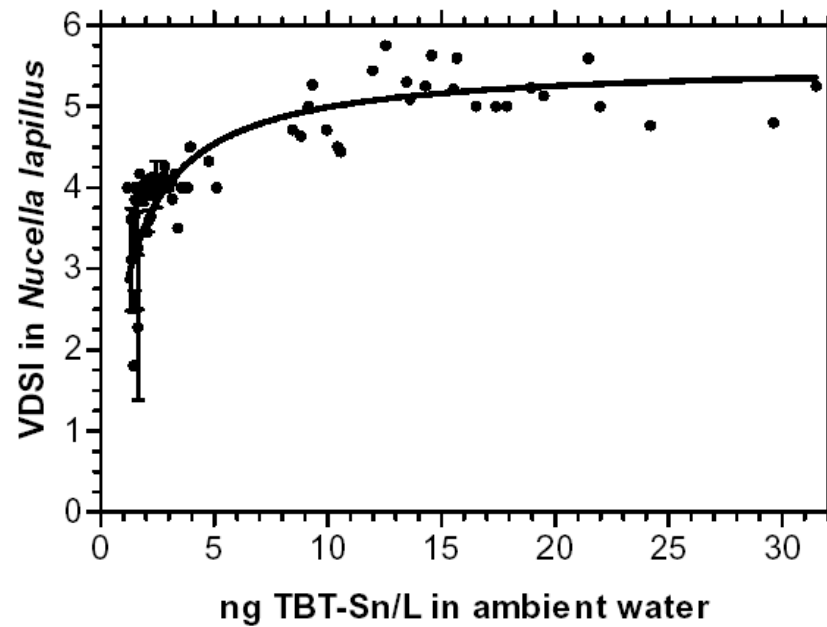


Fig. 2 *Nucella lapillus*. Relationship between aqueous TBT concentrations and imposex intensities:  $y = (5.54 x) / (1.12 + x)$ ;  $n = 151$  population samples from 81 stations;  $r = 0.688$ ;  $p < 0.0005$ .

One of the most important lessons to be learned from the "TBT story" and its effects in mollusks is that EDCs may impact different levels of biological integrations from molecules to communities affecting also the survival of populations in the field. Furthermore, the case history of TBT provides evidence for vertebrate-type steroids playing an important functional role in a number of invertebrate groups, including prosobranchs.

## Alternative Insecticides: Insect Growth Regulators

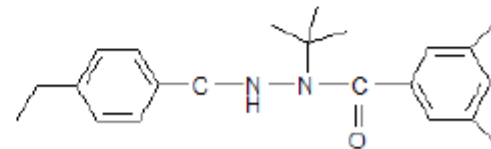
- “Nature and the pesticide industry apparently have decided that the best way to poison an animal is through its nervous system”
- most insecticides are nerve poisons, so mammalian selectivity must arise from differences in pharmacokinetics or metabolism
- a better way to go: find chemicals that attack biological processes *unique to insects* = biorational design
  - insects must shed their skin periodically to grow = molting
  - insects undergo metamorphosis between life stages
  - both processes are under strict endocrine control
  - 1. *juvenile hormone* (JH) titer in blood determines the next step in development; 2. *Ecdysone* (molting hormone) stimulates the molting process
- insecticides have been developed that mimic JH and ecdysone
- practical problem: agent must be present at the critical period to influence development (narrow window of susceptibility)

- Methoprene (Precor, Altosid) – first commercial *JH mimic* (Zoecon, 1978).  $LD_{50} = 34,600$ ,  $MSR = 1.7 \times 10^6$
- used as a mosquito larvicide (approved by WHO in drinking water for mosquito control); feed to livestock to control flies in manure; home control of fleas; control of stored product pests, mushroom pests. Used in Japan on silkworms to increase silk production.
- about 10 additional JH mimics have been commercialized



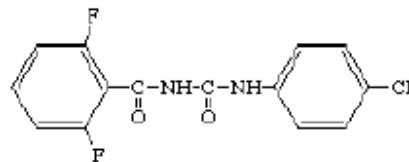
methoprene

- Tebufenozide (Mimic, Confirm) – first commercial *ecdysone agonist* (Rohm & Haas, 1991).  $LD_{50} > 5000$  mg/kg
- very effective against Lepidoptera and Colorado potato beetle – induces lethal premature molting
- first insecticide jointly registered by the EPA (USA) and the PMRA (Canada), 1996

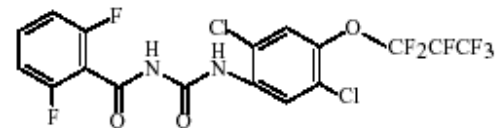


tebufenozide

- Diflubenzuron (Dimilin) – first commercial *chitin synthesis inhibitor* (Phillips, 1972).  $LD_{50} = 4640$  mg/kg.
- prevents insects from completing a molt by interfering with the synthesis of chitin, the main constituent of the integument. Not a direct action on chitin synthetase, but prevents the final step in activation of the enzyme. Slow acting.
- major uses: boll weevil on cotton, gypsy moth and other forest pests – used in Kitsilano in 1979 against gypsy moth



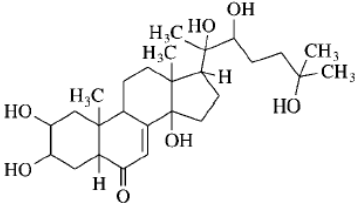
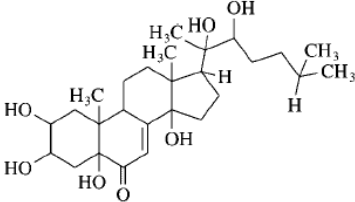
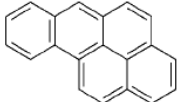
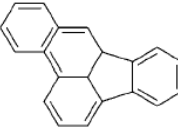
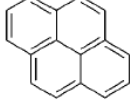
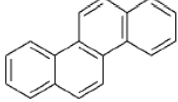
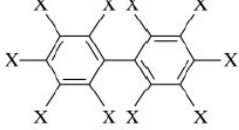
**diflubenauron**



**lufenuron**

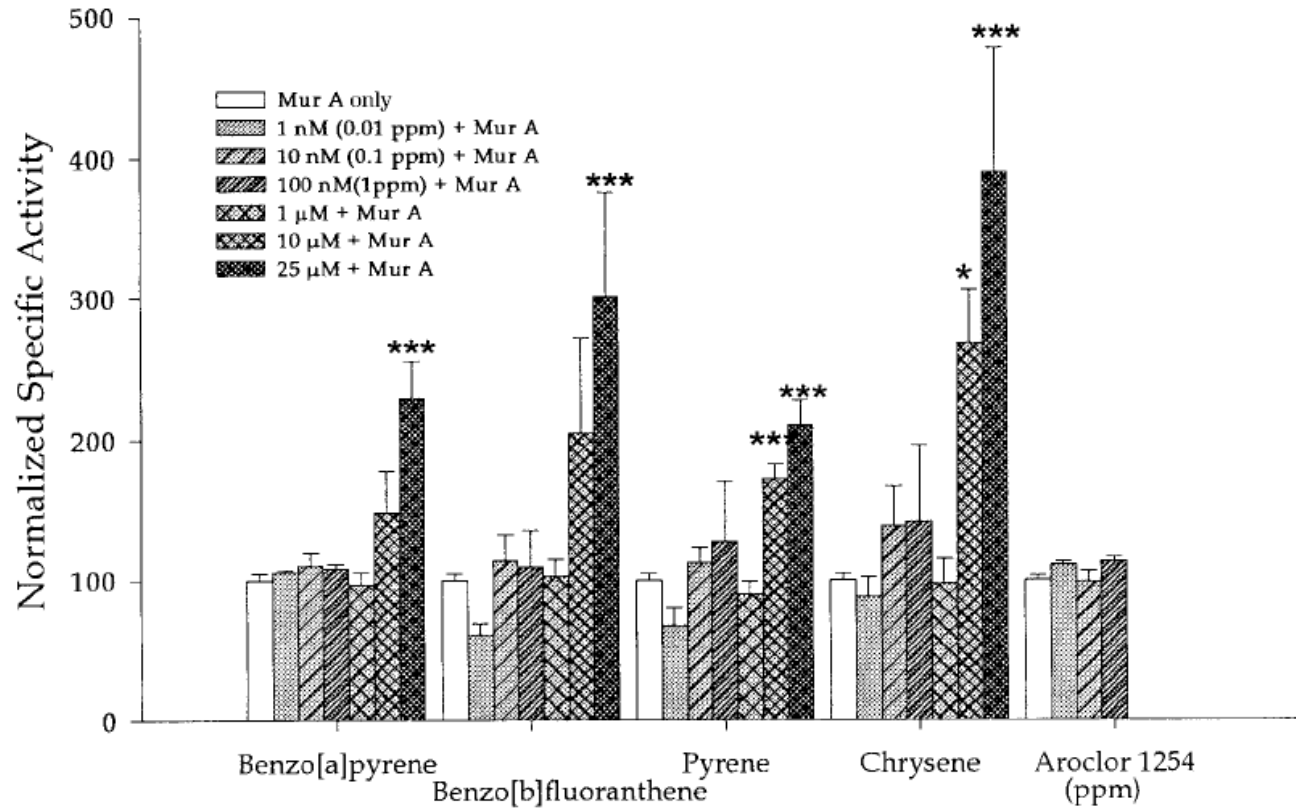
- Lufenuron (Program) – developed by Novartis in 1994. Oral IGR for flea control in dogs and cats – prevents egg development.
- $LD_{50} = 2000$  mg/kg; pet dose  $\sim 20$  mg/kg

# Polycyclic aromatic hydrocarbons can activate EcR:

| Compound                               | Chemical structure   | Steroid interactions   | References  |
|--|--|--|---|
| 20-OH Ecdysone<br>(20 HE)              |                          | Ecdysone agonist   | Cottam and Milner (1997)<br>Yao <i>et al.</i> (1993)  |
| Muristerone A<br>(MurA)                |                          | Ecdysone agonist   | Cottam and Milner (1997)<br>Yao <i>et al.</i> (1993)  |
| Benzo[ <i>a</i> ]pyrene<br>(BaP)       |                           | AhR agonist<br>Anti-estrogen<br>Anti-androgen<br>Not estrogenic in fish      | Hankinson (1995)<br>Tran <i>et al.</i> (1996)<br>Chang and Liao (1987)<br>Thomas and Smith (1993)   |
| Benzo[ <i>b</i> ]fluoranthene<br>(BbF) |                           | AhR agonist  | Hankinson (1995)  |
| Pyrene                                 |                          | No AhR activation  | Poland and Knutson (1982)   |
| Chrysene                               |                         | AhR agonist<br>No estrogen receptor<br>(ER) interaction                      | Hankinson (1995)<br>Tran <i>et al.</i> (1996)   |
| Aroclor 1254                           | <br>X = H or Cl, 54% Cl | AhR agonist<br>Thyroid hormone<br>Not antiestrogen<br>Estrogenic (?) in fish | Hankinson (1995)<br>Jacobson and Jacobson (1996)<br>Goldey <i>et al.</i> (1995)<br>Krishnan and Safe (1993)<br>Flouriot <i>et al.</i> (1995)<br>Thomas and Smith (1993) |



## Polycyklické aromatické uhlovodíky mohou aktivovat EcR:



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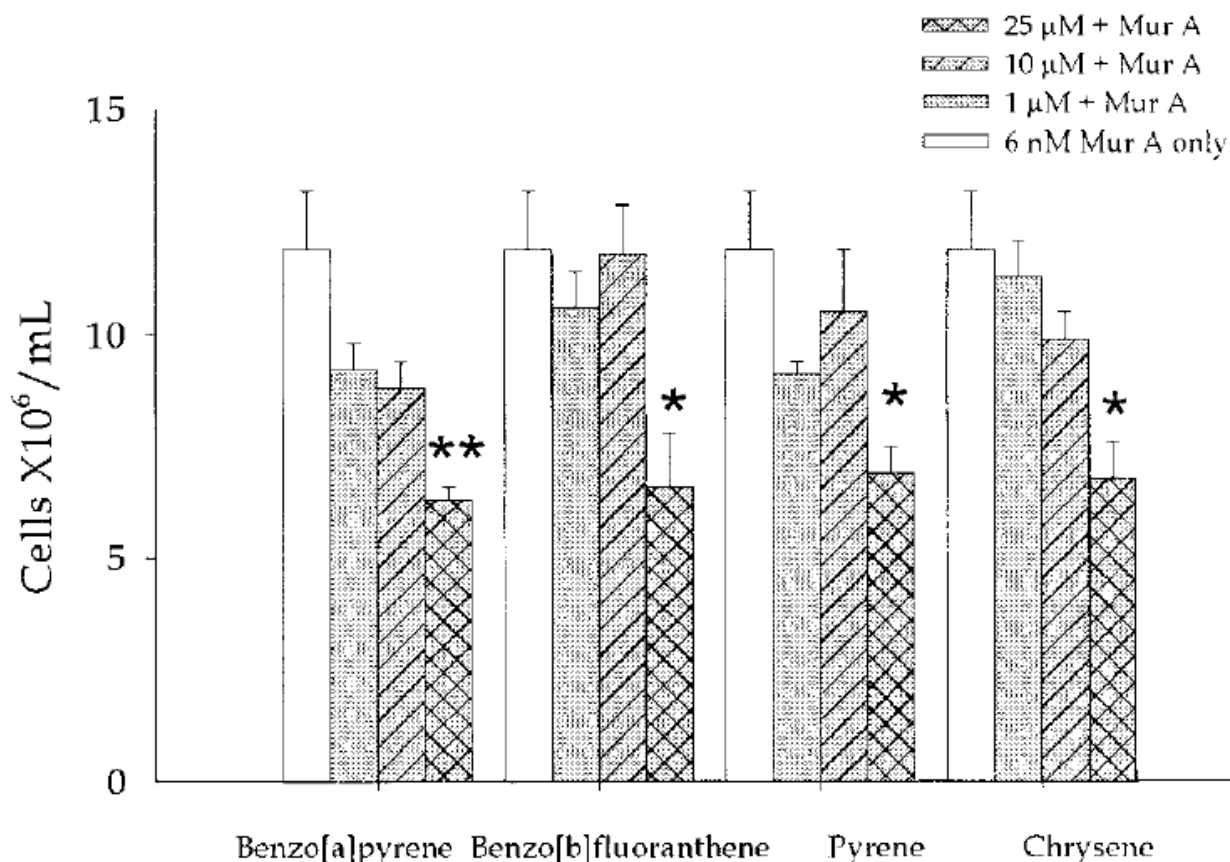


FIG. 4. Interaction of PAHs and Muristerone A in Cl.8+ cells. The four PAHs each enhance the differentiation response in Cl.8+ cells. This represents the average of three assays, each done in triplicate. Bars, SEM. \* $p < 0.05$ ; \*\* $p < 0.01$  different from 6 nM Muristerone A alone.

TABLE V

Representative Laboratory and Field Studies in Which Endocrine Disruption May Be Occurring in Insects<sup>a</sup>.

| Species                                     | Contaminant                           | Concentration (range), effects   | Lab/field     | Reference                       |
|---|---------------------------------------|--|---------------|---------------------------------|
| <i>Chironomus tentans</i> (all life stages) | 4-Nonylphenol                         | 12.5-200 µg/L; Reduced survival at high concentrations   | Lab           | Kahl <i>et al.</i> 1997         |
| <i>Macromia cingulata</i> (larvae)          | Tannery and paper pulp mill effluent  | 5, 10, 15, 20, 20%; Shortened time to first molt (tannery), arrested larval molting (paper pulp)                   | Lab           | Subramanian and Varadaraj 1993  |
| <i>Chironomus riparius</i> (larvae-adult)   | Phthalate esters                      | 100, 1000, 10000 mg/kg dw; No effects on survival, development or emergence  | Lab           | Brown <i>et al.</i> 1996        |
| <i>Chironomus</i> spp.                      | Industrial effluent containing metals | Ambient at site; Higher level of mentum deformities in exposed larvae in both field and lab conditions             | Field and Lab | Dickman and Rygiel 1996         |
| <i>Chironomus thummi</i> (larvae)           | Organic and inorganic pollutants      | Ambient at site; Prevalence of morphological deformities related to pollutants                                     | Field         | De Bisthoven <i>et al.</i> 1995 |
| <i>Chironomus</i> spp.                      | Possible exposure to pollutants       | Ambient at site; Deformed mouth parts, heavily pigmented head capsules, unusually thick head capsule and body wall | Field         | Hamilton and Saether 1971       |

<sup>a</sup>Modified from deFur *et al.* (1999).

TABLE III  
Representative Laboratory Studies in Which Endocrine Disruption May Be Occurring in Mollusks<sup>a</sup>.

| Species                             | Contaminant | Concentration (range),<br>Effects  | Lab/field | Reference                    |
|-------------------------------------|-------------|--|-----------|------------------------------|
| <i>Lymnaea stagnalis</i><br>(adult) | DDT, MCPA   | 50, 500 µg/L (DDT);<br>10, 100 mg/L (MCPA);<br>Fecundity alterations     | Lab       | Woin and<br>Bronmark<br>1992 |
| <i>Mytilus edulis</i><br>(adult)    | Cd          | 100 µg/L; Spawning<br>stimulation, inhibition of<br>gonadial development | Lab       | Kluytmans <i>et al.</i> 1988 |

<sup>a</sup>Modified from deFur *et al.* (1999).

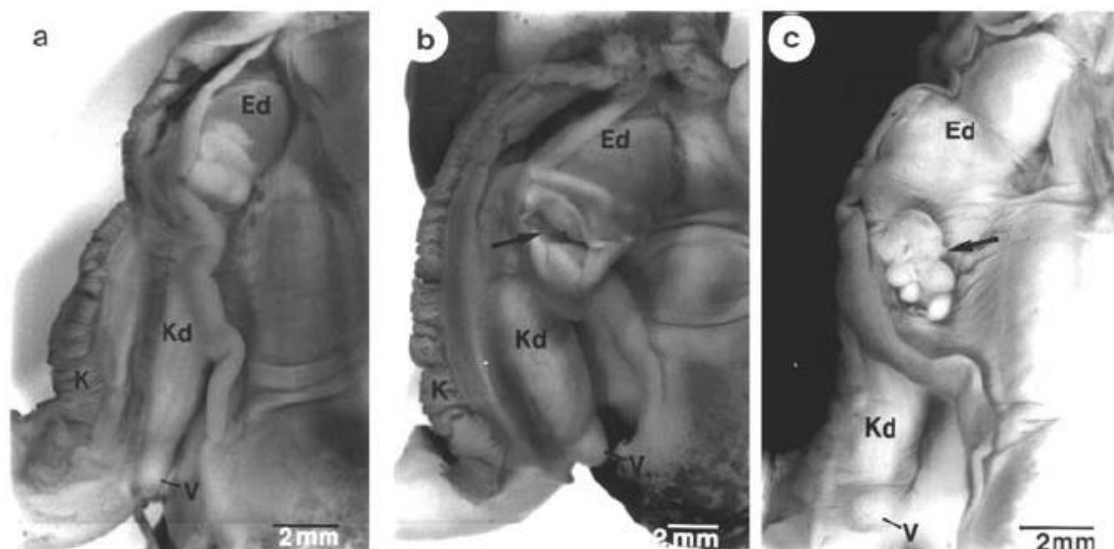
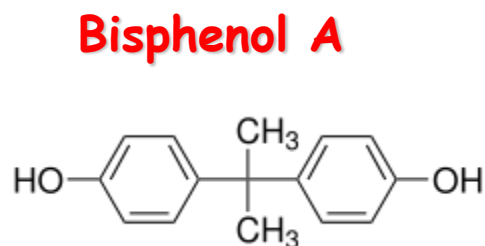


FIG. 3 *Marisa cornuarietis*. Photographs of a control female (a) and of BPA treated "superfemales" (b-c) with opened mantle cavities. In (b) and (c) a rupture in the wall of the pallial oviduct occurs (arrow) with an additional protrusion of the spawning mass in (c). Abbreviations: Ed, albumen gland; K, gill; Kd, capsule gland; V, vagina.

## Závěr

- Chemical signaling systems and their basic mechanisms in the animal kingdom exhibit a considerable degree of conservatism (McLachlan, 2001). Consequently, invertebrate endocrine function should be affected by identical or similar compounds as vertebrates (deFur *et al.*, 1999; Pinder and Pottinger, 1999).
- Highly effective EDCs have been intentionally developed for the purpose of pest control to interfere with hormonal systems of insects. Such endocrine-mediating properties can be assumed as not being unique for the IGRs or this group of arthropods but rather reflect the fact that much less research has been undertaken for other invertebrate groups than insects.
- ED in invertebrates found far less attention than in vertebrates in the past, probably because their hormonal systems are poorly understood favoring investigations with vertebrates and especially fish as systematic groups for ecotoxicological research and routine analyses many scientists feel familiar with.
- Little work has been done on endocrine disruption in invertebrates from the field (with the exception of the investigation of the imposex phenomenon in marine gastropods). The overwhelming majority of laboratory-based studies (56 reports) focuses on mollusks (17 publications), crustaceans (15 cases) and insects (12 reports), thus continuing main tendencies in the pre-1999 literature (Fig. 4a, b).