Development of effect based screening assays for the detection of marine toxins, using a genomics approach

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Why effect based assays? EU Regulations I

- Because we had good experience with the DR-CALUX screening for dioxins and PCBs (Friday Dr. Hädrich)
- We began with the hormones: Directive 96/23/EC: banns the use of Group A substances (list of compounds: targeted analytical analysis),
 - Stilbenes, derivatives, salts and esters
 - Antithyreogene compounds
 - Steroids
 - Resorcyclic Acid Lactones (including zeranol)
 - B-agonists

GENINGEN

• Others, as mentioned in the Annex of Regulation EC 37/2010



However,.... EU regulations II

Directive 96/23/EC mentions hormonal action and refers to Directive 96/22/EC

Directive 96/22/EC: Prohibits all substances having hormonal action

Regulations EC 178/2002 and EC 882/2004: oblige the member states to identify emerging risks and use validated and accredited methods for control analysis



How to obey to all these laws ?

The only way is bioactivity screening combined with chemical analytical confirmation and identification using validated and accredited methods for both

Or...to get rid of the laws. But would that be safe?





Bioactivity measurements for hormones

Transcriptional Activation (TA) bioassays (yeast or mammalian cell based)

Detect all compounds (structures) that are able to activate the receptor, e.g. the estrogen, androgen, progesterone or glucocorticoid receptor. As the main mode of action of these hormones is by activating their receptor, these TA bioassays fulfil Directive 96/22/EC that prohibits all substances having hormonal action, and they are also able to detect new designer steroids and new risks.

Moreover, they are:

- Sensitive and specific
- Quick, simple and robust
- Applicable to urine, feed and preparations

WAGENINGEN<mark>UR</mark>

Development of a yeast estrogen bioassay



The yeast estrogen bioassay





Bovee et al., *Gene* **325** (2004) 187-200 Bovee et al., *JSBMB* **91** (2004) 99-109

Similarly we developed a yeast androgen bioassay and yeast corticoid bioassay



Bovee et al., *ABC* **389** (2007) 1549-1558 Bovee et al., *ABC* **401** (2011) 873-882



I am not going to show you that:

SERMs and SARMs show their specific responses in these yeast hormone bioassays too

Both the yeast estrogen and androgen bioassay were fully validated for both the screening of feed and calf urine samples (according to Directive 2002/657/EC and accredited ISO 17025)

The yeast estrogen bioassay performed well in an interlaboratory ring test with calf urine samples

Was shown a cheap alternative for real practise: estrogen bioassay screening calf urine samples vs GC-MS analysis



Bovee et al., *JSBMB* **118** (2010) 85-92 Bovee et al., *ACA* **529** (2005) 57-64 Bovee et al., *FAC* **23** (2006) 556-568 Bovee et al., *ACA* **637** (2009) 225-234 Bovee et al., *ACA* **637** (2009) 265-272 Nielen et al., *FAC* **23** (2006) 1123-1131

I am also not going to show you that:

A 'natural' herbal supplement for prostate problems, causing gynaecomastia in a 67 year old man, was screened with the yeast estrogen bioassay and that it turned out that the supplement contained DES (Geldrop Hospital, The Netherlands) (This morning Dr. Weigel)



The yeast androgen bioassays specifically indicated the antiandrogenic potential of the printing ink compound 2isopropylthioxanthone (ITX), which was confirmed in vivo (rat) (food packaging)



Was validated by Waternet/Waterproef Laboratorium in The Netherlands for screening estrogens in water samples



Tooriaans et al., *FAC* **27** (2010) 917-925 Peijnenburg et al., *TiV* **24** (2010) 1619-1628 Nguyen et al., *TiV* **25** (2011) 2003-2009

And I am also not going to show you that:

- The yeast estrogen and androgen bioassay are successfully used at Ghent University for the screening of food supplements (including bio-directed identification)
- Both the yeast estrogen and androgen bioassay are used as the department of Food Chemistry, WUR for screening and bioassay-directed identification of active compounds in soy and licorice root



Both bioassays validated and used for calf urine and feed at the National Veterinary Research Institute, Poland

Both bioassays used for the screening of calf urine samples at the University of Veterinary Medicine, Turin, Italy



Becue et al., *ABC* **339** (2011) 1031-1039 Simons et al., *ABC* **401** (2011) 305-313 Simons et al., *J. Agric. Food Chem.* **59** (2011) 6748-6758 Divari et al., *FAC* **27** (2010) 1123-1131

Nor am I going to show you that:



- The yeast estrogen is successfully used at GSF, National Research Centre for Environment and Health, Neuherberg, Germany (Sediments)
- The yeast (gluco)corticoid assay was successfully used at Utrecht University-IRAS (hydroxylated PCBs)
- The yeast estrogen bioassay was successfully used at Utrecht University-IRAS (seawater contaminated with oil)

Is used at 1) Oregon State University, USA; 2) Amherst College, Chemistry Department, USA; 3) McGill University, Montreal Canada; 4) Chinese Academy of Science, China; 5) KFRI, Korea; 6) CRI, Bangkok, Thailand; 7) TU Dresden, Germany; 8) VITO, Belgium



Levy et al., *Environ. Sci. Pollut. Res. Int.* **18** (2011) 99-110 Antunes Fernandes et al., *Toxicology Letters* **206** (2011) 185-165 Vrabie et al., *Environ. Toxicol. Chem.* **29** (2010) 1529-1536

Nor that....



RIKILT: The yeast androgen assay was shown to of added value in a study with dietary supplements: a comparison with a liquid chromatography tandem mass spectrometry (LC-MS/MS) method (this morning Dr. Weigel)



RIKILT: Showed an added value by the identification of anabolic steroids and derivatives in supplements, using bioassay-guided fractionation, UHPLC/TOMMS analysis and accurate mass database searching





Rijk et al., *ACA* **637** (2009) 305-314 Peters et al., *ACA* **664** (2010) 77-88

But I am going to show you:



Approach of effect screening will work for marine toxins as well



Introduction

Food poisoning

- Bacteria
- Viruses
- Toxins



• Marine toxins

- ASP Amnesic Shellfish Poisc
- PSP Paralytic Shellfish Poison
- NSP Neurotoxic Shellfish Poison
- DSP Diarrheic Shellfish Poison



Introduction

Rats and mice are used for detecting marine toxins in mussels

- By feeding
- By injection



- From 2015 these methods will be forbidden (however not for testing production areas)
- LC-MS/MS method is available (for lipophilic toxins)
 - Expensive
 - Not high-throughput
 - Not developed to detect all marine toxins and unable to detect unknown toxins



I Development of a dedicated array for the detection of marine toxins. From genomics, via a dedicated tube array, back to the development of a specific bioassay?

Gene Selection (RIKILT): Caco-2 exposed to dinophysis toxin-1 (DTX1), azaspiracid 1 (AZA1) and okadaic acid (OA)

• Whole genome microarray (Service XS)

• Data analysed for selection marker genes



Gene selection (Caco-2: OA, DTX1 and AZA1)





Marine toxins – Genomics - Exposure of human Caco-2 cells

Genes specifically up- or down-regulated DTX1, AZA1 or OA were selected and used to develop a dedicated array (BioCop).

				INT_AVR vs	Intensity rank on
Gene	EXPRESSION GROUP	Spiked AZA1 AVR	Spiked DTX1 AVR	median	1 to 100
C21orf129	AZA and DTX1 down; BI_Mus_0	-1.76	-1.545	17.8	79
THEC	AZA and DTX1 down; BI_Mus_0	-1.55	-0.95	2.5	43
C3orf57	Uhiquely down by AZA1	-1.75	-0.12	4.7	56
NPPB	Spec down by DTX1	-0.07	-1.82	52.3	90
MT1H	Uhiquely up by AZA1, but also by higher DTX1 doses	2.155	0.23	22.3	82
MT1G	Uniquely up by AZA1, but also by higher DTX1 doses	2.375	0.23	16.4	78
CDKNIC	Uhiquely up by DTX1	-0.42	27	8.4	67
VASN	Uniquely up by DTX1	-0.06	3.1	4.6	55
MAFB	Uhiquely up by DTX1	0.09	3.495	1.7	34
RGS16	Uniquely up by DTX1	0.3	279	1.7	34
LOC387763	DTX1_>4x_AZA1_1,4_to_1.7_up	0.6	3.515	1.0	22
TUBB3	DTX1_>4x_AZA1_1,4_to_1.7_up	0.715	2.245	23.4	83
CEACAM1	AZA1 AND DTX1 UP	2.135	1.505	22.0	82
TNS4	AZA1 AND DTX1 UP	266	1.795	1.3	28
DDIT4	AZA1 AND DTX1 UP	2.4	1.435	18.7	80
TMCC1	Exp_Maria_AZA1_2,8up_DTX_max_1,3up	1.155	0.14	1.9	37
MT1F	Exp_Maria_AZA1_2,8up_DTX_max_1,3up	1.355	-0.5	11.0	72
TRIB3	Exp_Maria_AZA1_2,8up_DTX_max_1,3up	1.11	0.06	4.7	56
OSR2	Exp_Maria_AZA1_2,5down_DTX_max_1,4down	-0.845	0.81	2.8	46
AK091132	Exp_Maria_AZA1_2,5down_DTX_max_1,4down	-1.505	-0.175	3.4	50
GAPDH	Control	0.13	-0.14	318.9	99
TMEM179B	Control	-0.04	0.04	3.5	50



The newly developed Array Tube (Alere)

Transcriptomics assay on Clondiag AT-Platform >> ArrayTube (AT) Platform



single tube format based on conventional laboratory vials (Clondiag is nowadays Alere)

microtube



+ microarray



protein (HLA) array

- custom manufacturing of protein/peptide or nucleic acid based arrays
- array size of 2mm x 2mm with up to 300 features
- arrays including reaction control spots



easy processing with

no evaporation
uniform wettability
small volumes
optimal processing

standard lab equipment

through small surface area

oligonucleotide array



Array Tube Marine Toxins

Expose Caco-2 cells to standards or extracts (2-24h) and isolate the mRNA

 Synthesize the corresponding cDNA with a primerset of the 20 selected marker genes and 2 control genes (GAPDH and TMEM) (primerset 1)

Linear amplification with biotin labelled primerset 2 (in presence of competitor primer set 3, complementary to primers in set 1 and with an aminolink, in order to prevent an exponential amplification)

Hybridization on the Array Tube (staining was performed with a streptavidin coupled with a peroxidase and the addition of tetramethylbenzidin. The peroxidise activity catalyses the conversion of TMB and the dark blue precipitate product is measured by an array tube reader)

Measuring and data management (software) to get the results



Marine toxins – Development of a dedicated low-density microarray for genomic profiling





Bovee et al., ES&T 45 (2011) 8965-8973

Expression profile of the GAPDH control gene in the newly developed Array Tube





Expression profile of the CEACAM1 and TUBB3 marker genes in the Array Tube





Expression profile of the DDIT4 marker gene in the Array Tube





Bovee et al., ES&T 45 (2011) 8965-8973

Newly developed Array Tube for the detection of marine toxins

Thanks to the BioCop EU project (Prof. Elliott, QUB), Alere and the help from the group of Prof. Dr. Naegeli (Zürich)

It works, but....not all of the selected genes gave responses (7/20)

Labour intensive

Not cheap

Still takes about 3 days (mouse assay 1 day)



Alternatives – <u>Bioassays</u> or Molecular beacons or Multiplex qRT-PCR



- 1 Synthesize the -2.5 kb 5' from the ATG start of the DDIT gene (P) and combine with a marker (e.g. Luc or GFP) in a suited vector
- 2 E.g. Transfect the Caco-2 cell line with the DDITpromoter-Luc vector
- 3 Select clones on luciferase expression upon exposure to 1 nM AZA1



Alternatives – Bioassays or <u>Molecular beacons</u> or Multiplex qRT-PCR



A schematic illustration showing the concept of dual FRET molecular beacons. Hybridisation of donor and acceptor molecular beacons to adjacent regions on the same mRNA target results in FRET between donor and acceptor fluorophores upon donor excitation. By detecting FRET signal, fluorescence signals due to probe/target binding can be readily distinguished from that due to molecular beacon degradation and non-specific interactions [Santangelo et al., *Nucleic Acids Research* **32** (2004) 1-9].





Alternatives – Bioassays or Molecular beacons or Multiplex qRT-PCR



- 1 Take the Caco-2 cell
- 2 Expose cells and isolate mRNA

- 3 Perform qPCR on selected marker genes, e.g. DDIT4, CEACAM1, TUBB3, TRIB3 and OSR2



Gene Selection for qRT-PCR

- Based on the first exposure
- 22 genes were selected for the development of the Array Tube (DTX1, AZA1, OA)
- The complementary commercially designed Primers were ordered at Qiagen
- All tested individually in a SYBR green qRT-PCR (all worked well)
- 2nd exposure Caco-2 to yessotoxin (YTX) and pectenotoxin (PTX)
 - Whole genome microarray
 - Data analysed for selection of marker

 All tested individually in a SYBR green qRT-PCR

Qiagen – Primers and Probe design

TMEM179B -> TTTACTCCAACCTACAC
 RGS16 -> CCGCCTTCCCCACCAC
 DDIT4 -> TGTGTTTGTTGTTGTT
 NPPB -> CACCACGAAGCCCCAA

TMEM179B used as reference gene



RIKILT – single PCRs used to attribute the Probes

gene	dye	Excitation(nm)	Emission nm)
TMEM179B	-> Cy5	≻643	667
■RGS16	-> Texas Red	≻596	615
DDIT4	-> Hex	≻535	553
■NPPB	-> 6Fam	≻494	518



Results- "Singleplex" qRT-PCR (Biorad Sybr Green) & Multiplex qRT-PCR (Qiagen Quantifast Multiplex)







Able to detect ADTX, AZA, PTX and YTXAble to distinguish DTX and PTX



Future work

Developing the second multiplex qRT-PCR with the selected marker genes: CXCR4, EGR1 and TGFB2

• Detect OA

• Distinguish between AZA and YTX

	DTX	AZA	ΡΤΧ	OA	ΥΤΧ
CXCR4	1	?	?	\uparrow	?
EGR1	?	?	?	↑	?
TGFB2	↓	(\downarrow)	?	L L	?

TMEM179B used as reference gene



Future work

 Testing with blank and incurred mussel extracts (including positive in mouse bioassay, but negative by MS)

- Optimizing the method
 - Shorter exposure time
 - Faster RNA isolation kit
- Validation study
- Expand to other marine toxins (e.g. STX and SPX)
- Compare outcomes with a method being developed by a PhD student: using murine embryotic stem cell derived beating cardiomyocytes







