The Danish Pesticide Leaching Assessment Programme

Monitoring results May 1999-June 2008

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Preface

In 1998, the Danish Parliament initiated the Danish Pesticide Leaching Assessment Programme (PLAP), an intensive monitoring programme aimed at evaluating the leaching risk of pesticides under field conditions. The Danish Government funded the first phase of the programme from 1998 to 2001, while the Ministry of the Environment and the Ministry of Food, Agriculture and Fisheries funded a prolongation from 2002 to 2009.

The work was conducted by the Geological Survey of Denmark and Greenland (GEUS), the Faculty of Agricultural Sciences (DJF) at Aarhus University and the National Environmental Research Institute (NERI) Aarhus University under the direction of a management group comprising Jeanne Kjær (GEUS), Walter Brüsch (GEUS), Lis Wollesen de Jonge (DJF), Preben Olsen (DJF), Ruth Grant (NERI) and Steen Marcher (Danish Environmental Protection Agency).

This report presents the results for the period May 1999–June 2008. Results covering part of the period (May 1999–June 2007 have been reported previously (Kjær *et al.*, 2002, Kjær *et al.*, 2003, Kjær *et al.*, 2004, Kjær *et al.*, 2005c, Kjær *et al.*, 2007, and Kjær *et al.*, 2008). The present report should therefore be seen as a continuation of previous reports with the main focus on the leaching risk of pesticides applied during 2006.

The report was prepared jointly by Jeanne Kjær, Walter Brüsch, Vibeke Ernsten, Annette E. Rosenbom, Per Nyegaard, and Lasse Gudmundsson (all GEUS), Preben Olsen, and Finn Plauborg (DJF), and Ruth Grant (NERI). While all authors contributed to the whole report, the aspects for which authors were mainly responsible are as follows:

- Pesticide and bromide leaching: Walter Brüsch, Preben Olsen, and Jeanne Kjær.
- Soil water dynamics and water balances: Annette E. Rosenbom, Finn Plauborg, and Ruth Grant.
- Pesticide analysis quality assurance: Vibeke Ernsten.

Jeanne Kjær June 2009

Summary

In 1998, the Danish Parliament initiated the Pesticide Leaching Assessment Programme (PLAP), an intensive monitoring programme aimed at evaluating the leaching risk of pesticides under field conditions. The objective of the PLAP is to improve the scientific foundation for decision-making in the Danish registration procedures for pesticides. The specific aim is to analyse whether pesticides applied in accordance with current regulations leach to groundwater in unacceptable concentrations. The programme presently evaluates the leaching risk of 40 pesticides and 27 metabolites at five agricultural sites ranging in size from 1.1 to 2.4 ha. The results so far show that:

- Of the 40 pesticides applied, ten pesticides or their degradation products (clopyralid, chlormequat, desmedipham, florasulam, linuron, metsulfuron-methyl, thiamethoxam, tribenuronmethyl, iodosulfuron-methyl and triasulfuron) did not leach during the 1999-2008 monitoring period.
- The monitoring data indicate pronounced leaching of 12 of the applied pesticides and three of their degradation products. Thus ethofumesate, bentazone, propyzamide, tebuconazole, glyphosate and its degradation product AMPA, metamitron and its degradation product metamitron-desamino, azoxystrobin and its degradation product CyPM, as well as the degradation products of metribuzine, terbuthylazine, pirimicarb and rimsulfuron, picolinafen, leached through the root zone to 1 m below ground surface (hereafter m b.g.s.) in average concentrations exceeding 0.1 µg/l. Except for the degradation products of metribuzine, rimsulfuron and terbuthylazine, leaching was mainly confined to the depth of 1 m, where pesticides were frequently detected in the samples from suction cups and drainage systems.
- The monitoring data also indicate leaching of an additional 18 pesticides, in low concentrations, however. Thus, although the concentrations exceeded 0.1 µg/l in several samples, the average leaching concentrations at 1 m b.g.s. did not. Leaching was mainly confined to the depth of 1 m, where pesticides were detected in the samples from suction cups and drainage systems.

The PLAP initially evaluated the leaching risk at six agricultural sites representing a range of Danish soil and climate conditions. Monitoring at the Slaeggerup site was terminated on 1 July 2003, and results from that site are not included in the present report. For the monitoring results from this site see Kiær *et al.* (2004).

Pesticides were always applied in the maximum permitted dose. In order to describe water transport a bromide tracer was applied to the fields. Bromide and pesticide concentrations are measured monthly in both the unsaturated and the saturated zones, and weekly in the drainage water. This report covers the period May 1999—June 2008 and presents the monitoring results from the five agricultural sites presently monitored. The main focus is on evaluating the leaching risk of the pesticides applied during 2006.

Dansk sammendrag

I 1998 vedtog Folketinget at iværksætte projektet "Varslingssystem for udvaskning af pesticider til grundvandet" (VAP). VAP er et omfattende moniteringsprogram, der undersøger udvaskning af pesticider under reelle markforhold. Programmet har til formål at undersøge, om godkendte pesticider eller deres nedbrydningsprodukter – ved regelret brug – udvaskes til grundvandet i koncentrationer over grænseværdien for herigennem at udvide det videnskabelige grundlag for danske myndigheders (Miljøstyrelsens) procedurer for registrering af sprøjtemidler. Udvaskningrisikoen for 40 pesticider og 27 metabolitter således på fem marker af en størrelse på mellem 1,1 og 2,4 ha. De hidtidige resultater viser, at:

- Af de 40 pesticider, der er blevet udbragt, blev de ti eller nedbrydningsprodukter heraf (clopyralid, clormequat desmedipham, florasulam, linuron, metsulfuronmethyl, thiamethoxam, tribenuronmethyl og triasulfuron) ikke fundet udvasket i løbet af perioden 1999–2008.
- 12 af de udbragte stoffer eller nedbrydningsprodukter heraf gav anledning til en markant udvaskning. Ethofumesat, bentazon, propyzamid, tebuconazol, glyphosat, dettes nedbrydningsprodukt AMPA, metamitron, dettes nedbrydningsprodukt metamitron-desamino, azoxystrobin, dettes nedbrydningsprodukt CyPM, samt nedbrydningsprodukter fra henholdsvis metribuzin, picolinafen, terbuthylazin, pirimicarb og rimsulfuron blev udvasket fra rodzonen (1 meter under terræn, herefter m.u.t.) i gennemsnitskoncentrationer over 0,1 μg/l. På nær gældende metribuzins, rimsulfurons og terbutylazins nedbrydningsprodukter var udvaskningen primært begrænset til 1 m.u.t., hvor stofferne hyppigt blev fundet i prøver udtaget i sugeceller og dræn.
- Andre 18 stoffer gav anledning til udvaskning. Selv om flere af disse stoffer ofte blev fundet i koncentrationer over 0,1 μg/l, var der ikke tale om, at udvaskningen som årsmiddel oversteg 0,1 μg/l. Udvaskningen er primært begrænset til 1 m.u.t., hvor stofferne blev fundet i prøver udtaget i sugeceller og dræn

VAP-programmet omfattede oprindeligt seks marker placeret, så de repræsenterer forskellige typer geologi og tillige tager hensyn til de klimatiske variationer i Danmark, specielt hvad angår nedbørforhold. Monitering på den ene forsøgsmark (Slæggerup) stoppede 1. juli, 2003. Resultater fra denne mark er ikke inkluderet i denne rapport, men kan findes i Kjær *et al.* (2004). De anvendte pesticider bliver udbragt i maksimalt tilladte doser. Bromid anvendes som sporstof for at beskrive vandtransporten. Bromidog pesticidkoncentrationer bliver analyseret månedligt i prøver udtaget i den umættede og mættede zone og ugentligt i prøver af drænvand. I denne rapport præsenteres moniteringsresultaterne for de fem områder for perioden maj 1999-juni 2008 primært med fokus på pesticider udbragt i 2006. En del af stofferne har kun været inkluderet i moniteringsprogrammet i én udvaskningssæson, og for disse er det derfor for tidligt at konkludere noget endeligt.

1 Introduction

There is growing public concern in Denmark about pesticide contamination of our surface waters and groundwater. Pesticides and their degradation products have increasingly been detected in groundwater during the past decade and are now present in much of the Danish groundwater. Under the Danish National Groundwater Monitoring Programme (GRUMO) pesticides have so far been detected in 48% of all screens monitored and in 55% of the screens placed in the upper groundwater (Thorling. L. (red), 2009).

Due to the increasing detection of pesticides in groundwater over the past 10 years the desire has arisen to enhance the scientific foundation for the existing approval procedure for pesticides and to improve the present risk assessment tools. A main issue in this respect is that the EU assessment and hence also the Danish assessment of the risk of pesticide leaching to groundwater is largely based on data from modelling, laboratory or lysimeter studies. However, these types of data may not adequately describe the leaching that may occur under actual field conditions. Although models are widely used within the registration process, their validation requires further work, not least because of the limited availability of field data (Boesten, 2000). Moreover, laboratory and lysimeter studies do not include the spatial variability of the soil parameters (hydraulic, chemical, physical and microbiological soil properties) affecting pesticide leaching. This is of particular importance for silty and loamy soils, where preferential transport may have a major impact on pesticide leaching. In fact, various field studies suggest that considerable preferential transport of several pesticides occurs to a depth of 1 m under conditions comparable to those pertaining in Denmark (Kördel, 1997).

The inclusion of field studies, i.e. test plots exceeding 1 ha, in risk assessment of pesticide leaching to groundwater is considered an important improvement in risk assessment procedures. For example, the US Environmental Protection Agency (US-EPA) has included field-scale studies in its risk assessments since 1987. Pesticides that may potentially leach to the groundwater are required to be included in field studies as part of the registration procedure. The US-EPA has therefore conducted field studies on more than 50 pesticides (US Environmental Protection Agency, 1998). A similar concept has also been adopted within the European Union (EU), where Directive 91/414/EEC, Annexe VI (Council Directive 97/57/EC of 22 September 1997) enables field study results to be included in the risk assessments.

1.1 Objective

In 1998, the Danish Government initiated the Pesticide Leaching Assessment Programme (PLAP), an intensive monitoring programme aimed at evaluating the leaching risk of pesticides under field conditions. The PLAP is intended to serve as an early warning system providing decision-makers with advance warning if approved pesticides leach in unacceptable concentrations. The programme focuses on pesticides used in arable farming and monitors leaching at five agricultural test sites representative of Danish conditions.

The objective of the PLAP is to improve the scientific foundation for decision-making in the Danish registration and approval procedures for pesticides, enabling field studies to be included in risk assessment of selected pesticides. The specific aim is to analyse whether pesticides applied in accordance with current regulations leach at levels exceeding the maximum allowable concentration of $0.1 \,\mu g/l$.

1.2 Structure of the PLAP

The pesticides included in the PLAP were selected by the Danish Environmental Protection Agency on the basis of expert judgement. At present, 39 pesticides and several of their degradation products are included in the PLAP. All the compounds analysed are listed in Appendix 1.



Figure 1. Location of the PLAP sites Tylstrup, Jyndevad, Silstrup, Estrup and Faardrup. Monitoring at Slaeggerup was terminated on 1 July 2003.

Soil type and climatic conditions are considered to be some of the most important parameters controlling pesticide leaching. The PLAP initially encompassed six test sites representative of the dominant soil types and the climatic conditions in Denmark (Figure 1). Monitoring at the Slaeggerup site was terminated on 1 July 2003, and results from that site are not included in the present report. For the monitoring results from this site see Kjær et al. (2003). The groundwater table at all the sites is shallow, thereby enabling pesticide leaching to groundwater to be rapidly detected (Table 1). Cultivation of the PLAP sites is in line with conventional agricultural practice in the vicinity. The pesticides are applied in the maximum permitted dose and in the manner specified in the regulations. Hence any occurrence of pesticides or degradation products in the groundwater downstream of the sites can be related to the current approval conditions pertaining to the individual pesticides. The PLAP was initiated in autumn 1998. The five test sites encompassed by the present report were selected and established during 1999. Monitoring was initiated at Tylstrup, Jyndevad and Faardrup in 1999 and at Silstrup and Estrup in 2000 (Table 1).

Table 1. Characteristics of the five PLAP sites (modified from Lindhardt et al., 2001).

	Tylstrup	Jyndevad	Silstrup	Estrup	Faardrup
Location	Brønderslev	Tinglev	Thisted	Askov	Slagelse
Precipitation 1) (mm/y)	668	858	866	862	558
Pot. evapotransp. ¹⁾ (mm/y)	552	555	564	543	585
W x L (m)	70 x 166	135 x 184	91 x 185	105 x 120	150 x 160
Area (ha)	1.1	2.4	1.7	1.3	2.3
Tile drain	No	No	Yes	Yes	Yes
Monitoring initiated	May 1999	Sep 1999	Apr 2000	Apr 2000	Sep 1999
Geological characteristics					
– Deposited by	Saltwater	Meltwater	Glacier	Glacier/meltwater	Glacier
Sediment typeDGU symbol	Fine sand YS	Coarse sand TS	Clayey till ML	Clayey till ML	Clayey till ML
Depth to the calcareous matrix (m b.g.s.)Depth to the reduced matrix (m b.g.s.)	6 >12	5–9 10–12	1.3 5	1-4 ²⁾ >5 ²⁾	1.5 4.2
– Max. fracture depth ³⁾ (m)	_	_	4	>6.5	8
Fracture intensity 3–4 m depth (fractures/m)Ks in C horizon (m/s)	- 2.0·10 ⁻⁵	- 1.3·10 ⁻⁴	<1 3.4·10 ⁻⁶	11 8.0·10 ⁻⁸	4 7.2·10 ⁻⁶
	2.0 10	1.5.10	3.4.10	8.0.10	7.2.10
Topsoil characteristics – DK classification	JB2	JB1	JB7	JB5/6	JB5/6
- Classification	Loamy sand	Sand	Sandy clay loam/ sandy loam		Sandy loam
- Clay content (%)	6	5	18–26	10–20	14–15
- Silt content (%)	13	4	27	20–27	25
- Sand content (%)	78	88	8	50-65	57
– pH	4-4.5	5.6-6.2	6.7–7	6.5-7.8	6.4-6.6
– TOC (%)	2.0	1.8	2.2	1.7–7.3	1.4

¹⁾ Yearly normal based on a time series for the period 1961–90. The data refer to precipitation measured 1.5 m above ground.

2) Large variation within the field.

³⁾ Maximum fracture depth refers to the maximum fracture depth found in excavations and wells.

Site characterization and monitoring design are described in detail in Lindhardt et al. (2001). The present report presents the results of the monitoring period May 1999–June 2008. The main focus of this report is on the leaching risk of pesticides applied during 2006. For a detailed description of the earlier part of the monitoring period (May 1999–June 2006), see Kjær et al., (2002), Kjær et al. (2003), Kjær et al. (2004), Kjær et al. (2005c) and Kjær et al. (2007).

Under the PLAP the leaching risk of pesticides is evaluated on the basis of at least two years of monitoring data. For some pesticides the present report must be considered preliminary because they have been monitored for an insufficient length of time.

Hydrological modelling of the unsaturated zone at each PLAP site supports the monitoring data. The MACRO model (version 5.1), see Larsbo *et al.* (2005), was used to describe the soil water dynamics at each site during the entire monitoring period from May 1999 – June 2008. The five site models have been calibrated for the monitoring period May 1999 - June 2004 and validated for the monitoring period July 2004 - June 2008.

Scientifically valid methods of analysis are essential for the integrity of the PLAP. The field monitoring work has therefore been supported by intensive quality assurance entailing continuous evaluation of the analyses employed. The quality assurance methodology and results are presented in Section 7.

2 Pesticide leaching at Tylstrup

2.1 Materials and methods

2.1.1 Site description and monitoring design

Tylstrup is located in northern Jutland (Figure 1). The test field covers a cultivated area of 1.1 ha (70 x 166 m) and is practically flat, with windbreaks bordering the eastern and western sides. Based on two soil profiles dug in the buffer zone around the test field the soil was classified as a Humic Psammentic Dystrudept (Soil Survey Staff, 1999). The topsoil is characterized as loamy sand with 6% clay and 2.0% total organic carbon (Table 1). The aquifer material consists of about 20 meters of marine sand sediment deposited in the Yoldia Sea. The southern part is rather homogeneous, consisting entirely of fine-grained sand, whereas the northern part is more heterogeneous due to the intrusion of several silt and clay lenses (Lindhardt *et al.*, 2001). The overall direction of groundwater flow is towards the west (Figure 2). During the monitoring period the groundwater table was 2.6–4.5 m b.g.s. (Figure 3). A brief description of the sampling procedure is provided in Appendix 2. The monitoring design and test site are described in detail in Lindhardt *et al.* (2001), and the analysis methods in Kjær *et al.* (2002).

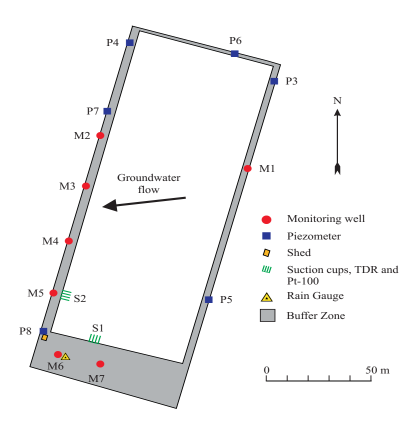


Figure 2.Overview of the Tylstrup test site. The innermost white area indicates the cultivated land, while the grey area indicates the surrounding buffer zone. The positions of the various installations are indicated, as is the direction of groundwater flow (by an arrow).

2.1.2 Agricultural management

Management practice during the 2006 and 2007 growing seasons is briefly summarized below and detailed in Appendix 3 (Table A3.1). For information about management practice during the previous monitoring periods, see Kjær *et al.* (2002, 2003, 2004, 2005c, 2007 & 2008).

Debris from a previous crop of maize was incorporated by rotary cultivation in November 2005 and in April 2006 the field was ploughed. A crop of spring barley (cv. Cabaret) was sown on 23 April 2006. Two weeks later the barley emerged. Weeds were sprayed with tribenuron-methyl on 6 June when the first node was detectable. The fungicide epoxiconazole was applied on 3 July at the end of heading. Due to a precipitation deficit the crop was irrigated twice with 29 mm on 19 June and 9 July. Harvest of the spring barley took place on 24 August yielding 55.6 hkg/ha of grain (85% dry matter), which was similar to the average for the year and location (Plantedirektoratet, 2006). The straw, 33.2 hkg/ha (100% dry matter), was shredded at harvest and incorporated the following day using a rotary cultivator.

On 26 August 2006 the field was ploughed and on the same day sown with winter rape seeds (cv. Lioness) dressed with the insecticide thiamethoxam, and sprayed with the herbicide clomazone (the latter, however, not included in the programme). Due to a heavy precipitation event shortly after sowing the surface of the soil was severely crusted. The crust impeded the emergence of the rape and on 6 September the field was resown (cv. Castille). The herbicides propyzamide and clopyralid were applied on 9 February and 27 March 2007, respectively. A single irrigation was applied using 30 mm on 8 June. The winter rape was harvested and straw shredded on 1 August, yielding 2.45 t/ha of seed (91% dry matter). Straw and stubble amounted to 6.47 t/ha (100% dry matter). The seed harvest was approximately 20% less than average for this soil type and year (Plantedirektoratet, 2008), most likely caused by the required late reseeding - see above.

The field was rotary cultivated twice on 3 August and 7 September 2007 in order to incorporate the rape residues. Ploughing on 12 September was followed by the sowing of winter wheat (cv. Smuggler) on 24 September, which emerged on 5 November. The herbicide pendimethalin was applied on 18 October at the stage of one unfurled leaf. The fungicide tebuconazol was used on 16 November at which time three leaves were unfurled. On 22 and 29 May 2008, 32 mm of irrigation was applied and on 5 and 13 June, 30 mm was applied. The fungicide azoxystrobin was used 17 June at the end of flowering, but not included in the monitoring. Harvest of winter wheat took place on 18 August, yielding 92.1 hkg/ha of grain (85% dry matter), which was 20% above the average for this soil type and year (Plantedirektoratet, 2008), and 18.5 hkg/ha of straw (100% dry matter).

2.1.3 Model setup and calibration

The numerical model MACRO (version 5.1) was applied to the Tylstrup site covering the soil profile to a depth of 5 m b.g.s., always including the groundwater table. The model was used to simulate water and bromide transport in the unsaturated zone during the full monitoring period May 1999–June 2008 and to establish an annual water balance.

Table 2. Annual water balance for Tylstrup (mm/y). Precipitation is corrected to soil surface according to the method of Allerup and Madsen (1979).

	Normal precipitation ²⁾	Precipitation	Irrigation	Actual evapotranspiration	Groundwater recharge 3)
1.5.99–30.6.99 1)	120	269	0	112	156
1.7.99-30.6.00	773	1073	33	498	608
1.7.00-30.6.01	773	914	75	487	502
1.7.01-30.6.02	773	906	80	570	416
1.7.02-30.6.03	773	918	23	502	439
1.7.03-30.6.04	773	758	0	472	287
1.7.04-30.6.05	773	854	57	477	434
1.7.05-30.6.06	773	725	67	488	304
1.7.06-30.6.07	773	1147	59	591	615
1.7.07-30.6.08	773	917	124	572	453

¹⁾ Accumulated for a two-month period.

Compared to Kjær *et al.* (2008), a year of validation was added to the MACRO-set-up for the Tylstrup site. The set-up was hereby calibrated for the monitoring period May 1999-June 2004 and validated for the monitoring period July 2004-June 2008. Daily time series of groundwater table measured in the piezometers located in the buffer zone, soil water content measured at three different depths (25, 60 and 110 cm b.g.s.) from the two profiles S1 and S2 (Figure 2) and the bromide concentration measured in the suction cups located 1 and 2 m b.g.s. were used in the calibration and validation process. Data acquisition, model set-up and results related to simulated bromide transport are described in Barlebo *et al.* (2007) and Rosenbom *et al.* (In prep.).

2.2 Results and discussion

2.2.1 Soil water dynamics and water balances

The model simulations were generally consistent with the observed data, thus indicating a good model description of the overall soil water dynamics in the unsaturated zone (Figure 3). The overall trends in soil water saturation were modelled successfully with the model capturing soil water dynamics at all depths (Figure 3C-E). The initial decrease in water saturation observed during the summer periods at 25, 60 and 110 cm b.g.s. was, however, less well captured. The dynamics of groundwater table were captured with some exceptions, but as with previous simulations the amplitude of the fluctuations was less well described (Figure 3B).

The resulting annual water balance is shown for each hydraulic year of the monitoring period (July–June) in Table 2. Precipitation and actual evapotranspiration of the most recent hydraulic year, July 2007–June 2008, lies within the interval observed since monitoring began at the site. The precipitation was characterized by being the fourth highest of the hydraulic years observed, with July, September and March being very wet, and May and June being very dry (Appendix 4). The very dry May and June resulted in the highest artificial irrigation applied to the field within the monitoring period and no simulated percolation 1 m b.g.s. in June (Figure 3A).

²⁾ Normal values based on time series for 1961–1990.

³⁾ Groundwater recharge is calculated as precipitation + irrigation - actual evapotranspiration.

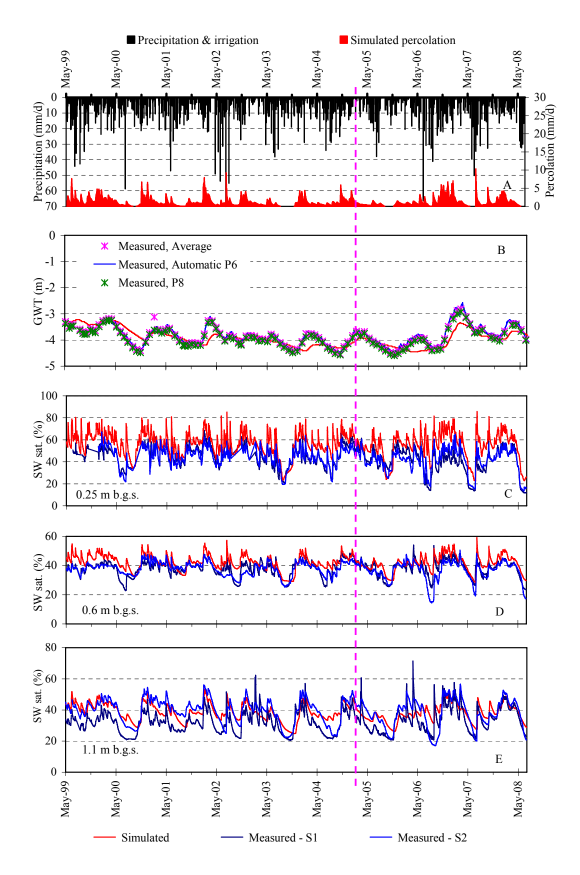


Figure 3. Soil water dynamics at Tylstrup: Measured precipitation, irrigation and simulated percolation 1 m b.g.s. (A), simulated and measured groundwater level GWT (B), and simulated and measured soil water saturation (SW sat.) at three different soil depths (C, D and E). The measured data in B derive from piezometers located in the buffer zone. The measured data in C, D and E derive from TDR probes installed at S1 and S2 (Figure 2). The broken vertical line indicates the beginning of the validation period (July 2004-June 2008).

2.2.2 Bromide leaching

Bromide has now been applied twice at Tylstrup. The bromide concentrations measured up to April 2003 (Figure 4 and Figure 5) relate to the bromide applied in May 1999, as described further in Kjær *et al.* (2003). Unsaturated transport of the bromide applied in March 2003 is evaluated in Barlebo *et al.* (2007) and Rosenbom *et al.* (In prep.)

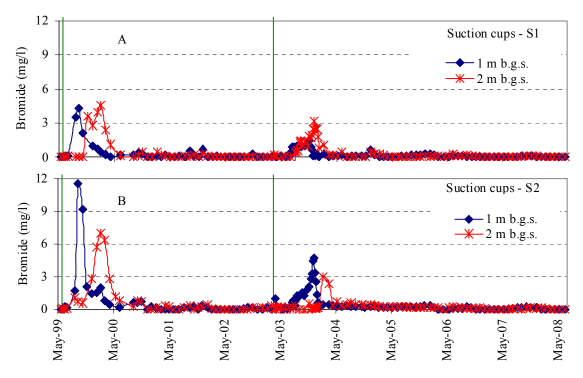


Figure 4. Measured bromide concentration in the unsaturated zone at Tylstrup. The measured data derive from suction cups installed 1 m b.g.s. and 2 m b.g.s. at locations S1 (A) and S2 (B) indicated in Figure 2. The green vertical lines indicate the dates of bromide applications.

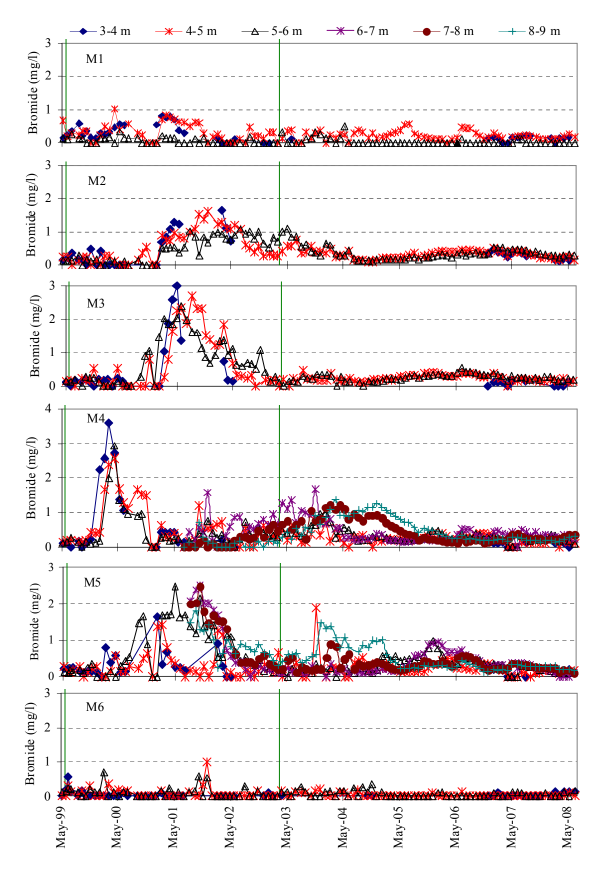


Figure 5. Bromide concentration in the groundwater at Tylstrup. The data derive from monitoring wells M1–M6. Screen depth is indicated in m b.g.s. The green vertical lines indicate the dates of bromide application.

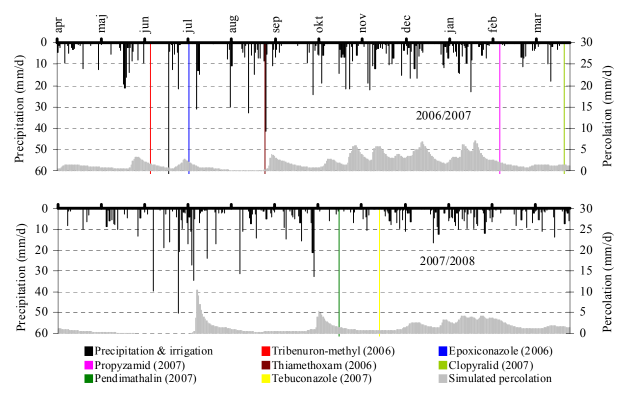


Figure 6. Pesticide application, precipitation and irrigation (primary axis) together with simulated percolation 1 m b.g.s. (secondary axis) at Tylstrup in 2006/2007 (upper) and 2007/2008 (lower).

2.2.3 Pesticide leaching

Monitoring at Tylstrup began in May 1999 and presently encompasses several pesticides and their degradation products, as shown in Table 3. Pesticide applications during the two most recent growing seasons are shown together with precipitation and simulated precipitation in Figure 6.

It should be noted that precipitation in Table 3 is corrected to soil surface according to Allerup and Madsen (1979), whereas percolation (1 m b.g.s.) refers to accumulated percolation as simulated with the MACRO model. It should also be noted that as mancozeb (applied here as Dithane DG), tribenuron-methyl (applied here as Express) and rimsulfuron (applied here as Titus) degrade rapidly, the risk of leaching risk is more associated with their respective degradation products, ETU, triazinamin-methyl, PPU and PPU-desamino. This is why the degradation products and not the parent compounds are monitored in the PLAP (Table 3). Pesticides applied later than April 2008 are not evaluated in this report and hence are not included in Table 3 and Figure 6.

The leaching risk of pesticides applied from 2000 to 2005 has been evaluated in Kjær *et al.* (2002, 2003, 2004, 2005c, 2007 & 2008). The leaching of metribuzin is further detailed in Kjær *et al.* (2005b).

Table 3. Pesticides analysed at Tylstrup with the products used shown in parentheses. Degradation products are in italics. Precipitation (prec.) and percolation (perc.) are accumulated from the date of first application until the end of monitoring. 1^{st} month perc. refers to accumulated percolation within the first month after the application. C_{mean} refers to average leachate concentration at 1 m b.g.s. the first year after application. The number of pesticide-positive samples is indicated in parentheses.

Crop and analysed pesticides	Application	End of	Prec.	Perc.	1 st month	C _{mean}
	date	monitoring	(mm)	(mm)	perc. (mm)	(µg/L)
Potatoes 1999						
Linuron (Afalon)	May 99	Jul 01	2550	1253	87	< 0.01 (0)
- $ETU^{(1)}$ (Dithane DG)	Jun 99	Oct 01	2381	1169	73	< 0.01(9)
Metribuzine (Sencor WG)	Jun 99	Jul 03	4223	2097	85	< 0.01 (3)
- metribuzine-diketo		Jul 08 [†]	8918	4235	85	0.05 - 0.36(659)
- metribuzine-desamino		Jul 03	4223	2097	85	< 0.02(0)
- metribuzine-desamino-diketo		Apr 08	8689	4192	85	0.14-0.97 (317)
Spring barley 2000						
Triasulfuron (Logran 20 WG)	May 00	Apr 03	2740	1283	13	< 0.02(0)
- triazinamin	-	-				< 0.02 (0)
Propiconazole (Tilt Top)	Jun 00	Jul 03	2948	1341	11	< 0.01 (0)
Fenpropimorph (Tilt Top)	Jun 00	Jul 03	2948	1341	11	< 0.01 (0)
- fenpropimorphic acid						< 0.02(0)
Pirimicarb (Pirimor G)	Jun 00	Apr 03	2622	1263	17	< 0.01 (0)
- pirimicarb-desmethyl						< 0.02 (0)
- pirimicarb-desmethyl-formamido						< 0.02 (0)
Winter rye 2001						
Pendimethalin (Stomp SC)	Nov 00	Apr 03	2271	1219	109	< 0.01 (0)
Triazinamin-methyl 2) (Express)	Nov 00	Apr 03	2271	1219	109	< 0.02 (0)
Propiconazole (Tilt Top)	May 01	Jul 03	2948	1341	11	< 0.01 (0)
Fenpropimorph (Tilt Top)	May 01	Jul 03	2948	1341	11	< 0.01 (0)
- fenpropimorphic acid						< 0.01 (0)
Winter rape 2002						
Clomazone (Command CS)	Sep 01	Jul 04	2534	1194	9	< 0.01 (0)
- propanamide-clomazone	-					< 0.02 (0)

Systematic chemical nomenclature for the analysed pesticides is given in Appendix 1.

However, rimsulfuron applied in 2004 continued to leach, and the results of this 2004 application are summarised below.

Rimsulfuron degrades rapidly in the soil, and the leaching risk is therefore more associated with the degradation products PPU and PPU-desamido. PPU has been found several times in suction cups situated 1 m and 2 m b.g.s. at both S1 and S2. The first detection of PPU occurred at S1, 10 months after the application (Figure 7B), whereafter PPU was found in 20 out of 27 analysed samples with concentrations ranging between 0.021 and 0.092 μ g/l. A little more than two years after application PPU was found at 1 m depth in S2 in several samples, concentrations ranging between 0.01 and 0.067 μ g/l (Figure 7D). Small concentrations of PPU were seen in both S1 and S2 at the end of the monitoring period, indicating that although leaching slowly decreased, it had not yet ceased. The decreased concentration observed in 2008 may be correlated to a small infiltration of water from the root zone at the start of 2008. Average concentrations did not exceed 0.1 μ g/l in any of the three years for either of the metabolites (Table 4). Minor leaching of PPU-desamido was also seen at Tylstrup, but the number of detections and concentration levels were low (Figure 7 and Appendix 5).

^{1]} Degradation product of mancozeb. The parent compound degrades too rapidly to be detected by monitoring.

²⁾ Degradation product of tribenuron-methyl. The parent compound degrades too rapidly to be detected by monitoring.

[†] Monitoring will continue during the following year. The values for prec. and perc. are accumulated up to July 2008.

Table 3 continued. Pesticides analysed at Tylstrup with the products used shown in parentheses. Degradation products are in italics. Precipitation (prec.) and percolation (perc.) are accumulated from date of first application until the end of monitoring. 1st month perc. refers to accumulated percolation within the first month after the application. C_{mean} refers to average leachate concentration at 1 m b.g.s. the first year after application. The number of pesticidepositive samples is indicated in parentheses.

Crop and analysed pesticides	Application	End of	Prec.	Perc.	1st month	C _{mean}
	date	monitoring	(mm)	(mm)	perc. (mm)	$(\mu g/l)$
Winter wheat 2003						
Bromoxynil (Oxitril CM)	Oct 02	Apr 05	2082	995	53	< 0.01 (0)
Ioxynil (Oxitril CM)	Oct 02	Apr 05	2082	995	53	< 0.01 (0)
Fluroxypyr (Starane 180)	May 03	Jul 05	1867	787	50	< 0.02 (0)
Flamprop-M-isopropyl (Barnon Plus 3) - Flamprop-M (free acid)	May 03	Jul 05	2635	1031	42	<0.01 (0)
Dimethoate (Perfekthion 500 S)	Jul 03	Jul 05	1629	722	14	< 0.01 (0)
Potatoes 2004						
-Fluazifop-P (free acid) 1) (Fusilade X-tra)	May 04	Jul 06	1754	704	16	< 0.01 (0)
- PPU ²⁾ (Titus)	Jun 04	Jul 08 [†]	3987	1856	13	<0.01 (66) 3)
- PPU- desamino ²⁾ (Titus)	Jun 04	Jul 08 [†]	3987	1856	13	<0.01 (28) 3)
Maize 2005						
Terbuthylazine (Inter-Terbutylazine)	May 05	Jul 07	2145	933	16	< 0.01 (0)
-desethyl-terbuthylazine	,					<0.01(2)
-2-hydroxy-terbuthylazine						< 0.01(1)
-desisopropyl-atrazine						<0.01 (18) 3)
-2-hydroxy-desethyl-terbuthylazine						< 0.01 (6)
Bentazone (Laddok TE)	Jun 05	Jul 07	2061	927	33	< 0.01(1)
-AIBA						< 0.01 (0)
Spring barley 2006						
-triazinamin-methyl ⁴⁾ (Express ST)	Jun 06	Jul 08	2349	1184	43	< 0.02 (0)
Epoxiconazole (Opus)	Jul 06	Jul 08	2233	1148	24	< 0.01(0)
Winter rape 2007						
Thiamethoxam (Cruiser RAPS) ⁵⁾	Aug 06	Apr 08	2030	1123	57	< 0.01 (0)
-CGA 322704		P				<0.02 (0)
Propyzamide (Kerb 500 SC)	Feb 07	Jul 08 [†]	1385	599	40	< 0.01 (0)
-RH-24644						< 0.01(0)
-RH-24580						< 0.01 (0)
-RH-24655						< 0.01 (0)
Clopyralid (Matrigon)	Mar 07	Jul 08 [†]	1302	539	24	< 0.01 (0)
Winter wheat 2008						
Pendimethalin (Stomp)	Oct 07	Jul 08 [†]	627	357	27	< 0.01 (0)
Tebuconazole (Folicur EC250)	Nov 07	Jul 08 [†]	591	332	46	<0.01(1)

PPU was found in a single groundwater sample (0.045 µg/l on 7 December 2005 in the monitoring screen of M4 located 4.4-5.4 m b.g.s.); PPU-desamido has not been seen in the groundwater (Table A5.1 in Appendix 5).

Systematic chemical nomenclature for the analysed pesticides is given in Appendix 1.

Degradation product of fluazifop-P-butyl. The parent compound degrades too rapidly to be detected by monitoring.

²⁾ Degradation product of rimsulfuron. The parent compound degrades too rapidly to be detected by monitoring.

³⁾ Leaching increased the second year after application (see Figure 7).
4) Degradation product of tribenuron-methyl. The parent compound degrades too rapidly to be detected by monitoring.

⁵⁾ Unlike the other pesticide applied via surface spray application, thiamethoxam was directly applied in the soil as the rape seeds (cv. Lioness) were dressed with thiamethoxam.

[†] Monitoring will continue during the following year. The values for prec. and perc. are accumulated up to July 2008.

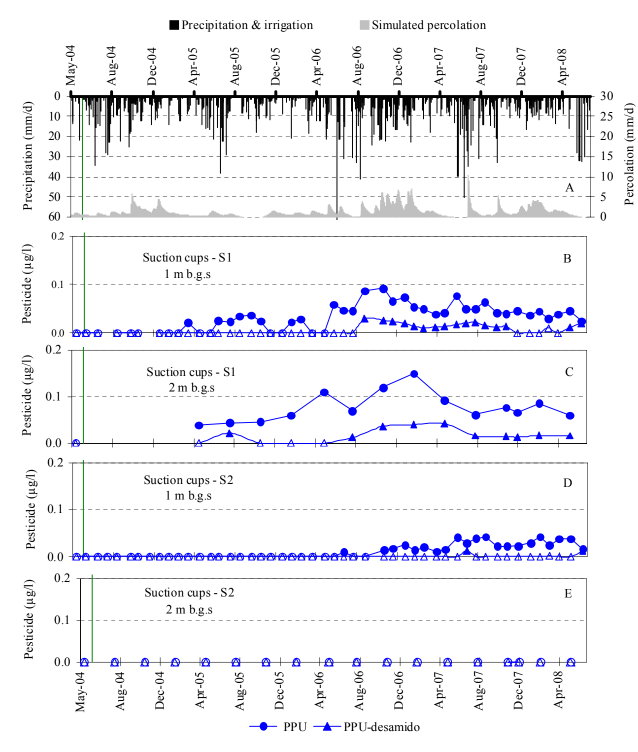


Figure 7. Precipitation, irrigation and simulated percolation 1 m b.g.s. (A) together with measured concentration of *PPU* and *PPU* –desamino (μ g/l) in suction cups installed on location S1 at 1m b.g.s. (B) and 2 m b.g.s. (C) and location S2 at 1m b.g.s. (D) and 2 m b.g.s. (D) at Tylstrup. The green vertical line indicates the date of pesticide application. Open circles and triangles indicate concentrations below the detection limit of 0.02 μ g/l.

Table 4. Percolation together with estimated average concentration (μ g/l) of PPU and PPU-desamido 1 m b.g.s. at Tyletrum

	Percolation	PPU		PPU-desamido		
	(mm/y)	Suction cup – S1	Suction cup – S2	Suction cup – S1	Suction cup – S2	
1.7.04–30.6.05	528	< 0.02	< 0.02	< 0.02	< 0.02	
1.7.05-30.6.06	257	0.02	< 0.02	< 0.02	< 0.02	
1.7.06-30.6.07	529	0.07	0.02-0.03	< 0.02	< 0.02	
1.7.07-30.6.08	514	0.04-0.04	0.03-0.03	0.01-0.01	<0.01-0.01	

When evaluating these results it should be noted that precipitation following the application of rimsulfuron (applied on 3 June 2004) amounted to 68 mm in May 2004 (20% higher than normal) and 51 mm in June 2004 (21% lower than normal). Precipitation and percolation following the application at Tylstrup were thus much lower than that observed at Jyndevad in 2003 where rimsulfuron was also applied. Finally, it should be noted that the concentration of PPU is likely to be underestimated by up to 28-47%. Results from the field-spiked samples thus indicating that PPU is unstable and tends to degrade further to PPU-desamido during storage and transport (see section 7.2.2.). Thus, the observed PPU-desamido probably derives from degradation in the sample during subsequent storage and transport rather than from degradation occurring in the soil. As a consequence, the concentration of PPU is likely to be underestimated, while that of PPU-desamido is likely to be overestimated.

In 2006 tribenuron-methyl and epoxiconazole were applied to spring barley in June. The parent compound tribenuron-methyl degrades rapidly and therefore only the metabolite triazineamine-methyl is monitored. However, no trace of the pesticide or the metabolite was found in water samples taken from groundwater or suction cups in the monitoring period 2006-2008 (Table 3).

The pesticides applied on winter rape 2007 and winter wheat 2008 and their metabolites (Table 3) have not yet been found in any of the analysed water samples. The final evaluation awaits an additional year of monitoring.

3 Pesticide leaching at Jyndevad

3.1 Materials and methods

3.1.1 Site description and monitoring design

Jyndevad is located in southern Jutland (Figure 1). The test site covers a cultivated area of 2.4 ha (135 x 184 m) and is practically flat. A windbreak borders the eastern side of the test site. The area has a shallow groundwater table ranging from 1 to 3 m b.g.s. (Figure 9B) The overall direction of groundwater flow is towards northwest (Figure 8). The soil can be classified as Arenic Eutrudept and Humic Psammentic Dystrudept (Soil Survey Staff, 1999) with coarse sand as the dominant texture class and topsoil containing 5% clay and 1.8% total organic carbon (Table 1). The geological description points to a rather homogeneous aquifer of meltwater sand, with local occurrences of thin clay and silt beds. A brief description of the sampling procedure is provided in Appendix 2. The monitoring design and test site are described in detail in Lindhardt et al. (2001) and the analysis methods in Kjær et al. (2002).

3.1.2 Agricultural management

Management practice during the 2006-2007 growing seasons is briefly summarized below and detailed in Appendix 3 (Table A3.2). For information about management practice during the previous monitoring periods see Kjær *et al.* (2002, 2003, 2004, 2005c, 2007 & 2008).

The field was limed on 13 March 2006 using 3.3 t/ha of dolomite chalk (92%), rotary-cultivated on 29 March and ploughed the subsequent day. Two weeks later spring barley (cv. Simba) was sown. The barley emerged on 21 April. The herbicide florasulam was sprayed on 26 May when the second node was detectable. On 8 June, at heading, the fungicide epoxiconazole was applied. Due to a dry growing season, the field was irrigated five times. An amount of 27, 30, and 27 mm was applied on 6, 12 and 26 June. On 2 and 7 July 30 mm was applied. Harvest of barley took place on 7 August yielding 60.5 hkg/ha of grain (85% dry matter) and 26.8 hkg/ha of straw, which was approximately 10 hkg/ha grain more than similar, irrigated soils in the region (Plantedirektoratet, 2006).

The field was ploughed on 20 September 2006 and sown with triticale (cv. Dinaro) the following day. Emergence of the triticale took place on 1 October. The herbicides mesosulfuron and iodosulfuron (Atlantis WG) were applied on 10 October, at the stage of three unfurled leaves. Due to the risk of lodging, chlormequat was applied on 13 April 2007 at the beginning of stem elongation. The crop was irrigated with 27 mm on 27 April. Fungicidal treatment using epoxiconazole was carried out on 7 May at the beginning of heading. A second irrigation, was conducted on 5 June, again using 27 mm. On 7 August the crop was harvested yielding 38.7 hkg/ha (85% dry matter) as well as 38.3 hkg/ha of straw (100% dry matter). Grain yields were low at 13 hkg/ha less than average for that soil type and year (Plantedirektoratet, 2006).

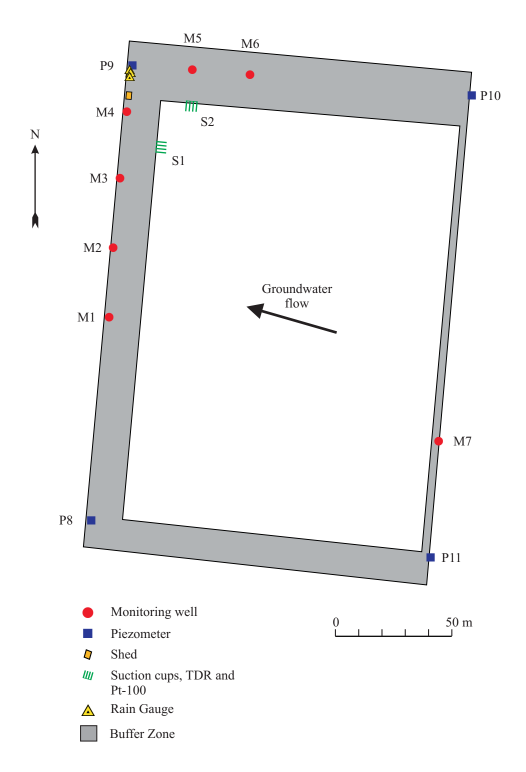


Figure 8. Overview of the Jyndevad test site. The innermost white area indicates the cultivated land, while the grey area indicates the surrounding buffer zone. The positions of the various installations are indicated, as is the direction of groundwater flow (by an arrow).

Having been sprayed with glyphosate (not included in the monitoring) on 13 September, the field was ploughed on 28 September and sown with winter wheat (cv. Ambitron) on 1 October 2007. The crop emerged on 12 October. The herbicide picolinafen was applied on 29 October and the fungicide tebuconazol on 3 December. In both instances the crop had two unfurled leaves. Due to the very dry growing season of 2008, the field was irrigated on seven occasions: 7 May (42 mm), 14 May (27 mm), 21 May (27 mm), 30 May (30 mm), 5 June (35 mm), 25 June (35 mm) and 8 July (30 mm). The fungicide azoxystrobin was applied on 11 June at full flowering. Harvest took place on 30 August, yielding 68.1 hkg/ha grain (85% dry matter) and 28.1 hkg/ha straw (100% dry matter), grain yield being similar to the average for this soil type that year (Plantedirektoratet, 2008).

3.1.3 Model set-up and calibration

The numerical model MACRO (version 5.1, Larsbo *et al.* (2005)) was applied to the Jyndevad site covering the soil profile to a depth of 5 m b.g.s., always including the groundwater table. The model was used to simulate water flow and bromide transport in the unsaturated zone during the full monitoring period July 1999–June 2008 and to establish an annual water balance.

Compared with the set-up in Kjær *et al.* (2008), a year of validation was added to the MACRO-set-up for the Jyndevad site. The set-up was hereby calibrated for the monitoring period May 1999-June 2004 and validated for the monitoring period July 2004-June 2008. For this purpose, the following time series have been used: the groundwater table measured in the piezometers located in the buffer zone, soil water content measured at three different depths (25, 60, and 110 cm b.g.s.) from the two profiles S1 and S2 (location indicated at Figure 8), and the bromide concentration measured in the suction cups located 1 and 2 m b.g.s (Figure 11). Data acquisition, model set-up as well as results related to simulated bromide transport are described in Barlebo *et al.* (2007) and Rosenbom *et al.* (In prep.).

3.2 Results and discussion

3.2.1 Soil water dynamics and water balances

The model simulations were generally consistent with the observed data, thus indicating a good model description of the overall soil water dynamics in the unsaturated zone (Figure 9). The dynamics of the simulated groundwater table was well described with MACRO 5.1 (Figure 9B). As noted earlier (Kjær *et al.*, 2008), the model had some difficulty in capturing the degree of soil water saturation 1.1 m b.g.s. – a difference that is still apparent (Figure 9E). As in the Tylstrup scenario, the decrease in water saturation observed during the summer periods at 25 and 60 cm b.g.s. was less well captured.

Table 5. Annual water balance for Jyndevad (mm/yr). Precipitation is corrected to the soil surface according to the method of Allerup and Madsen (1979).

	Normal Precipitation 1)	Precipitation	Irrigation	Actual Evapotranspiration	Groundwater recharge ²⁾
1.7.99–30.6.00	995	1073	29	500	602
1.7.00-30.6.01	995	810	0	461	349
1.7.01-30.6.02	995	1204	81	545	740
1.7.02-30.6.03	995	991	51	415	627
1.7.03-30.6.04	995	937	27	432	531
1.7.04-30.6.05	995	1218	87	578	727
1.7.05-30.6.06	995	857	117	490	484
1.7.06-30.6.07	995	1304	114	571	847
1.7.07-30.6.08	995	1023	196	613	605

¹⁾ Normal values based on time series for 1961–1990.

The resulting water balance for Jyndevad for the nine monitoring periods is shown in Table 5. Compared with the previous eight years, the latest hydraulic year July 2007-June 2008 was characterised by having the third highest precipitation, the highest simulated actual evapotranspiration and the highest irrigation values. Precipitation in the latest hydraulic year was characterized by July and March being very wet, and October, April and May being very dry (Appendix 4). Due to this precipitation pattern, continuous percolation 1 m b.g.s. was simulated throughout the year.

²⁾ Groundwater recharge is calculated as precipitation + irrigation - actual evapotranspiration.

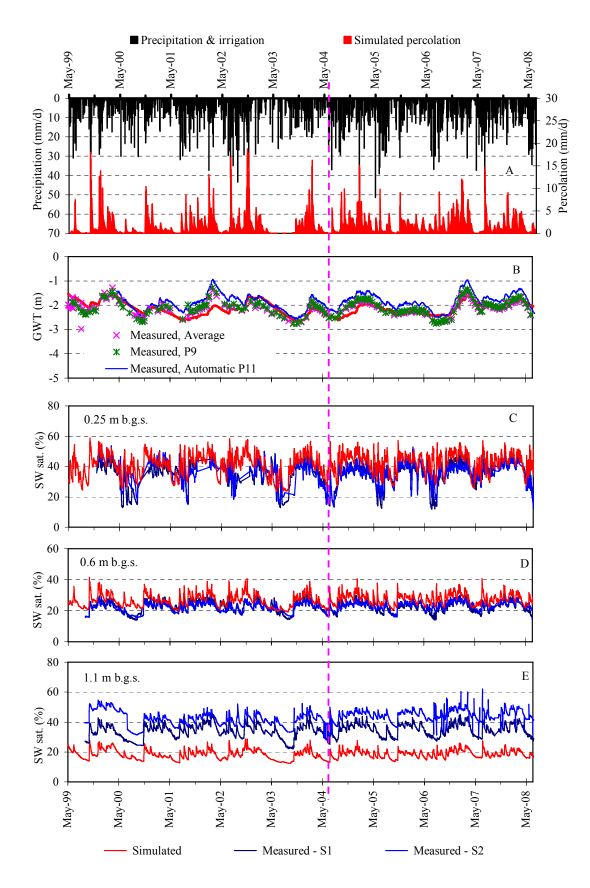


Figure 9. Soil water dynamics at Jyndevad: Measured precipitation, irrigation and simulated percolation 1 m b.g.s. (A), simulated and measured groundwater level (B), and simulated and measured soil water saturation (SW sat.) at three different soil depths (C, D and E). The measured data in B derive from piezometers located in the buffer zone. The measured data in C, D and E derive from TDR probes installed at S1 and S2 (see Figure 8). The broken vertical line indicates the beginning of the validation period (July 2004- June 2008).

3.2.2 Bromide leaching

Bromide has now been applied twice at Jyndevad. The bromide concentrations measured up to April 2003 (Figure 10 and Figure 11) relate to the bromide applied in autumn 1999, as described further in Kjær *et al.* (2003). Leaching of the bromide applied in March 2003 is evaluated in Rosenbom *et al.* (In prep.).

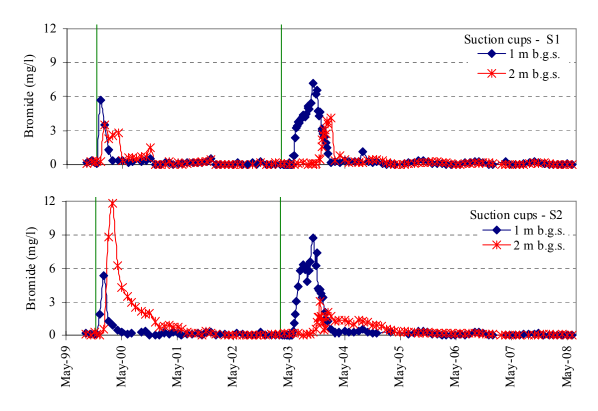


Figure 10. Bromide concentration in the unsaturated zone at Jyndevad. The measured data derive from suction cups installed (A) 1 m b.g.s. and (B) 2 m b.g.s. at locations S1 and S2 (Figure 8). The green vertical lines indicate the dates of bromide applications.

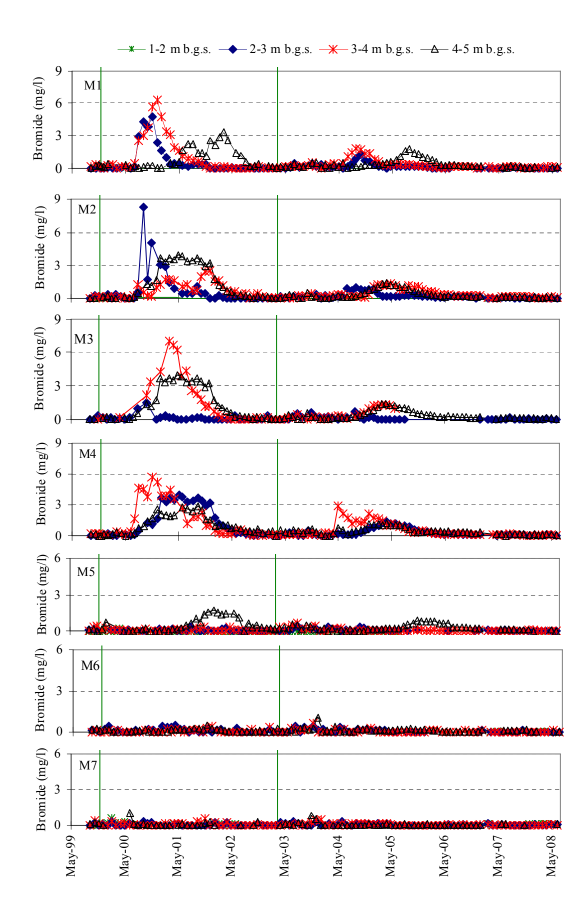


Figure 11. Bromide concentration in the groundwater at Jyndevad. The data derive from monitoring wells M1–M7. Screen depth is indicated in m b.g.s. The green vertical lines indicate the dates of bromide applications.

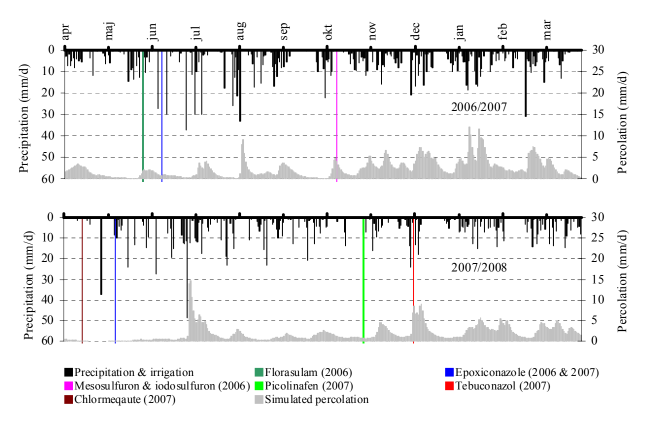


Figure 12. Pesticide application, precipitation and irrigation (primary axis) together with simulated percolation 1 m b.g.s. (secondary axis) at Jyndevad in 2006/2007 (upper) and 2007/2008 (lower).

3.2.3 Pesticide leaching

Monitoring at Jyndevad began in September 1999 and presently encompasses several pesticides and their degradation products, as indicated in Table 6. Pesticide application during the two most recent growing seasons is shown together with precipitation and simulated precipitation in Figure 12. It should be noted that precipitation is corrected to the soil surface according to Allerup and Madsen (1979), whereas percolation (1 m b.g.s.) refers to accumulated percolation as simulated with the MACRO model. It should also be noted that as tribenuron-methyl (applied here as Express), pyridate (applied here as Lido) and rimsulfuron (applied here as Titus) degrade rapidly, the leaching risk is more associated with their respective degradation products, triazinamin-methyl, PHCP, PPU and PPU-desamido. For the same reasons it is the degradation products and not the parent compounds that are monitored in the PLAP (Table 6). Pesticides applied later than April 2008 are not evaluated in this report and hence are not included in Table 6 and Figure 12.

Table 6. Pesticides analysed at Jyndevad with the product used shown in parentheses. Degradation products are in italics. Precipitation (prec.) and percolation (perc.) are accumulated from date of first application until end of monitoring. 1st month perc. refers to accumulated percolation within the first month after application. C_{mean} refers to average leachate concentration 1 m b.g.s the first year after application. The number of pesticide-positive samples is indicated in parentheses.

Crop and analysed pesticides	Application	End of	Prec.	Perc.	1 st month	C _{mean}
	date	monitoring	(mm)	(mm)	perc. (mm)	(μg/l)
Winter rye 2000						
Glyphosate (Roundup 2000)	Sep 99	Apr 02	2759	1607	139	< 0.01 (0)
- AMPA		-				< 0.01 (3)
Triazinamin-methyl 1) (Express)	Nov 99	Apr 02	2534	1451	86	< 0.02 (0)
Propiconazole (Tilt Top)	Apr 00	Jul 02	2301	1061	3	< 0.01 (0)
Fenpropimorph (Tilt Top)	Apr 00	Apr 02	2015	1029	3	< 0.01(2)
- fenpropimorphic acid	-					< 0.01 (0)
Maize 2001						
Terbuthylazine (Lido 410 SC)	May 01	Apr 04	3118	1809	4	< 0.01 (0)
- desethyl-terbuhylazine	May 01	Apr 07	6742	3826	4	<0.01-0.02 (42)
PHCP ²⁾ (Lido 410 SC)	May 01	Jul 03	2413	1366	4	<0.02(0)
Potatoes 2002						
- PPU (Titus) ³⁾	May 02	Jul 08 [†]	7085	3873	11	$0.06^{4} - 0.13$ (294)
- PPU-desamido (Titus)		Jul 08 [†]	7085	3873	11	0.01–0.03 (137)
Spring barley 2003						*****
MCPA (Metaxon)	Jun 03	Jul 05	2340	1233	0	< 0.01 (0)
- 4-chlor,2-methylphenol	Juli 03	Jul 03	2340	1233	U	<0.01 (0)
Dimethoate (Perfekthion 500 S)	Jun 03	Jul 05	2278	1232	1	<0.01 (0)
Pea 2004	54 11 05	our os	2270	1232	•	0.01 (0)
Bentazone (Basagran 480)	May 04	Jul 07	3888	2044	4	0.02-0.13 (30)
- AIBA	May 04	Jul 07	3000	2044	7	<0.01 (2)
Pendimethalin (Stomp SC)	May 04	Apr 07	3557	1996	4	<0.01 (2)
Pirimicarb (Pirimor G)	Jun 04	Apr 07	3493	1993	27	<0.01 (0)
- Pirimicarb-desmethyl	Juli 01	ripi o i	3 173	1775	27	<0.01 (1)
-Pirimicarb-desmethyl-						<0.02 (0)
formamido						10.02 (0)
- fluazifop-P(free acid) 5)	Jun 04	Jul 06	2395	1233	27	< 0.01 (0)
(Fusilade X-tra)						(1)
Winter wheat 2005						
Ioxynil (Oxitril CM)	Oct 04	Apr 07	2955	1791	81	< 0.01 (0)
Bromoxynil (Oxitril CM)	Oct 04	Apr 07	2955	1791	81	< 0.01(0)
Amidosulfuron (Gratil 75 WG)	Apr 05	Jul 07	1070	515	33	< 0.01(3)
Fluroxypyr (Starane 180 S)	May 05	Jul 07	2683	1360	37	< 0.02 (0)
Azoxystrobin (Amistar)	May 05	Apr 07	2274	1283	49	< 0.01 (0)
- CyPM	-	•				< 0.02 (0)
Spring barley 2006						
Florasulam (Primus)	May 06	Jul 08	2779	1487	34	< 0.01 (0)
- florasulam-desmethyl	J					< 0.03 (0)
Epoxiconazole (Opus)	Jun 06	Jul 08 [†]	2728	1465	34	< 0.01 (0)
Triticale 2007						
Mesosulfuron-methyl (Atlantis	Oct 06	Jul 08 [†]	2206	1291	73	< 0.01 (0)
WG)				, -	, -	< 0.01 (0)
- mesosulfuron						()
Chlormequat (Cycocel 750)	Apr 07	Jul 08 [†]	1548	689	1	< 0.01 (0)
Epoxiconazole (Opus)	May 07	Jul 08 [†]	1222	648	6	< 0.01 (0)
Winter wheat 2008						
Picolinafen (Pico 750 WG)	Oct 07	Jul 08 [†]	836	462	55	< 0.01 (0)
- CL153815						<0.01(0)
Tebuconazole (Folicur EC 250)	Dec 07	Jul 08 [†]	709	402	97	< 0.01 (0)

Systematic chemical nomenclature for the analysed pesticides is given in Appendix 1.

1) Degradation product of tribenuron-methyl. The parent compound degrades too rapidly to be detected by monitoring.

2) Degradation product of pyridate. The parent compound degrades too rapidly to be detected by monitoring.

3) Degradation product of rimsulfuron. The parent compound degrades too rapidly to be detected by monitoring.

Degratation product of rimsultaron. The parent compound degrates too rapidly to be detected by monitoring.

4) Leaching increased the second year after application (see Figure 13).

5) Degradation product of fluazifop-P-butyl. The parent compound degrades too rapidly to be detected by monitoring.

† Monitoring will continue for an additional year. The values for prec. and perc. are accumulated up to July 2008.

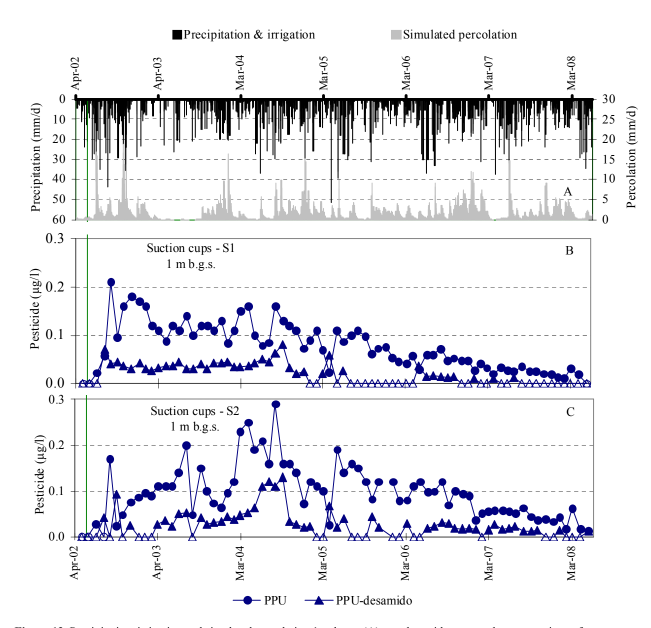


Figure 13. Precipitation, irrigation and simulated percolation 1 m b.g.s. (A) together with measured concentrations of *PPU* and *PPU-desamido* (μ g/l) in suction cups installed 1 m b.g.s. at location S1 (B) and S2 (C) at Jyndevad. The green vertical line indicates the date of pesticide application. Open circles and triangles indicate concentrations below the detection limit of 0.02 μ g/l.

The leaching risk of pesticides applied during 2000, 2001, 2002, 2003, 2004, and 2005 has been evaluated in Kjær et al. (2002, 2003, 2004, 2005c, 2007 & 2008). Since PPU and PPU-desamido (metabolites of rimsulfuron applied in 2003) were still included in the current monitoring period, the results of these applications are, however, summarised below. For a detailed description of the leaching pattern, including primary data and climatic conditions characterising the monitoring periods, reference is made to Kjær et al. (2005c).

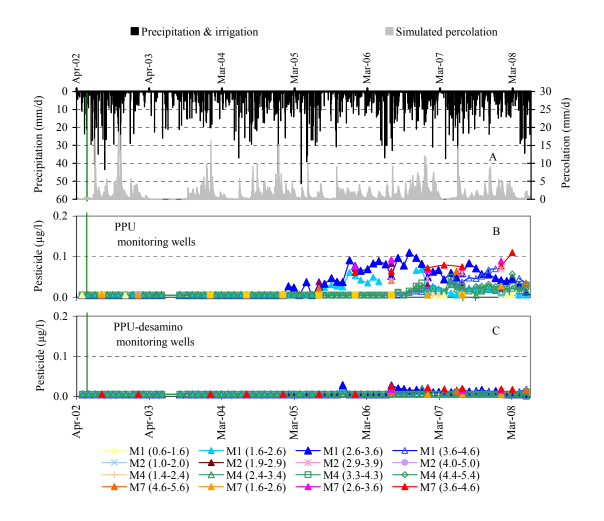


Figure 14. Precipitation, irrigation and simulated percolation 1 m b.g.s. (A) together with measured concentrations $(\mu g/l)$ in downstream (M1, M2, M4) and upstream monitoring wells (M7) of *PPU* (B) and *PPU-desamido* (C) at Jyndevad. The numbers in parentheses indicate the depths of the analysed screens. The green vertical line indicates the date of pesticide application. Open circles and triangles indicate concentrations below the detection limit of 0.02 $\mu g/l$.

Two degradation products of rimsulfuron, PPU and PPU-desamido, were detected at 1 m depth in the suction cups at S1 and S2 (Figure 13). Both compounds were characterized by continuous leaching over a long period of time. Although the concentration decreased during the last monitoring year, PPU was still detected towards the end of 2008, i.e. six years after application. Average yearly concentrations of PPU reaching 0.1 µg/l were seen as long as three years after application (Figure 13 and Table 7). With an overall travel time of about four years PPU also reached the downstream monitoring screens (Figure 14). Elevated concentrations of PPU were seen in M1 (sampled monthly), M2 (sampled half-yearly) and from 2006 in M4 (sampled monthly). PPU was also found in low concentrations in M7, which receives water from the upstream neighbouring field (Figure 9 and Figure 15). The PPU found in this well likely originated from the neighbouring upstream fields. This is because the tracer test suggested that water sampled in M7 had not infiltrated at the PLAP site, and because rimsulfuron was also applied at the neighbouring upstream area (Kjær et al. 2007).

Table 7. Percolation together with estimated average concentrations (μ g/l) of PPU and PPU-desamido 1 m b.g.s. at Jyndevad. Leached mass refers to the total mass (% of applied rimsulfuron) leached during the monitoring period 1 7 02–30 6 07

	Percolation	PP	U	PPU-desamido			
	(mm/y)	Suction cup – S1 Suction cup – S2 S		Suction cup – S1	Suction cup – S2		
1.7.02-30.6.03	706	0.13	0.06	0.03-0.04	0.02-0.03		
1.7.03-30.6.04	468	0.12	0.10	0.04	0.04		
1.7.04-30.6.05	759	0.10	0.14	0.03	0.05		
1.7.05-30.6.06	465	0.07	0.09	< 0.02	< 0.02		
1.7.06-30.6.07	815	0.05	0.08	0.02	0.02		
1.7.07-30.6.08	643	0.02	0.04	< 0.01-0.01	0.01		

¹⁾ Expressed as rimsulfuron equivalent.

In addition to the leaching observed at the PLAP site, this result thus indicates that leaching also occurred in the neighbouring upstream field. Furthermore, PPU-desamino was detected in monitoring wells, the number of detections and concentration level being lower than that of PPU-desamido, however (Figure 14C and Table A5.2 in Appendix 5).

Finally, it should be noted that the concentration of PPU is likely to be underestimated by up to 22-44% due to stability problems as described in Section 2.2.3 and section 7.2.2.

The leaching risk of the five pesticides applied during the 2006 growing season (florasulam and its metabolite florasulam-desmethyl, epoxiconazole, mesosulfuron-methyl and its metabolite mesosulfuron) was found to be negligible, as none of these compounds have yet been detected in any of the analysed samples.

The pesticides used on spring barley in 2007 have not yet been found in any of the analysed water samples. The final evaluation awaits an additional year of monitoring.

4 Pesticide leaching at Silstrup

4.1 Materials and methods

4.1.1 Site description and monitoring design

The test field at Silstrup is located south of Thisted in north-western Jutland (Figure 1). The cultivated area is 1.69 ha (91 x 185 m) and slopes gently 1–2° to the north (Figure 15). Based on two profiles excavated in the buffer zone bordering the field the soil was classified as Alfic Argiudoll and Typic Hapludoll (Soil Survey Staff, 1999). The topsoil content of clay in the two profiles was 18 and 26%, and the organic carbon content was 3.4 and 2.8%, respectively, Table 1. The geological description showed rather homogeneous clay till rich in chalk and chert, containing 20–35% clay, 20–40% silt and 20–40% sand. In some intervals the till was sandier, containing only 12–14% clay. Moreover, thin lenses of silt and sand were found in some of the wells. The gravel content was approx. 5%, but could be as high as 20%. A brief description of the sampling procedure is provided in Appendix 2. The monitoring design and test site are described in detail in Lindhardt *et al.* (2001) and the analysis methods in Kjær *et al.* (2002).

4.1.2 Agricultural management

Management practice during the 2006-2007 growing seasons is briefly summarized below and detailed in Appendix 3 (Table A3.3). For information about management practice during the previous monitoring periods see Kjær *et al.* (2002, 2003, 2004, 2005c, 2007 & 2008).

Within two hours after trailer hose application of pig slurry on 29 August, 2005, the field was ploughed and sown with winter rape (cv. Calypso) on 1 September. Before the emergence of the rape, the herbicide clomazone was sprayed onto the bare soil. On 17 November, when the rape had six unfurled leaves, the herbicide propyzamide was applied. On 20 April 2006, when the rape had nine visible internodes, the herbicide clopyralid was used. Pest control was performed on 5 May using alpha-cypermethrin. However, this substance was not included in the monitoring programme. The crop was fertilised with 23.8 t/ha pig slurry on 26 April 2006. On 24 July, when nearly all pods had reached the final size, the crop was windrowed. The crop was threshed on 7 August, yielding 37.3 hkg/ha of seeds (91% dry matter), a yield similar to the average on this type of soil within the region (Plantedirektoratet. 2006). All of the straw, 67.7 hkg/ha 100% dry matter, was shredded on the day of harvest. The field was ploughed on 20 September and two days later sown with winter wheat (cv. Skalmeje) as well as sprayed with the herbicide pendimethalin. Eight days later the wheat emerged. At the beginning of stem elongation, 13 June 2007, the plant growth inhibitor chlormequat and the herbicide iodosulfuron were applied. On 7 June, at the end of heading, the fungicide epoxiconazole was applied. The winter wheat was harvested on 24 August, yielding 100, 7 hkg/ha of grain and 40.8 hkg/ha straw – 85 and 100% of dry matter, respectively. The straw was shredded at harvest. Incorporation of the straw and stubble was done on 29 August using a disk harrow and the field was ploughed on 12 November. Grain yield was some 0.5 t/ha larger than on a similar soil for this year (Plantedirektoratet, 2008).

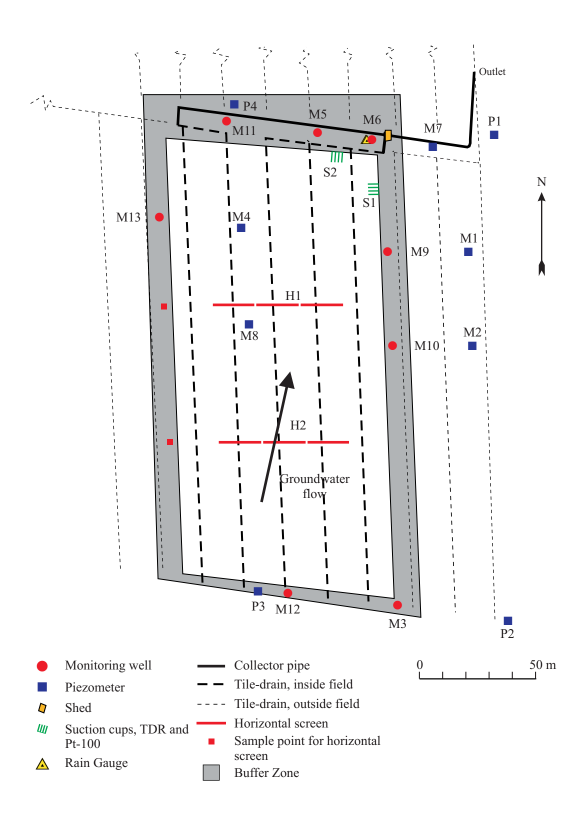


Figure 15. Overview of the Silstrup site. The innermost white area indicates the cultivated land, while the grey area indicates the surrounding buffer zone. The positions of the various installations are indicated, as is the direction of groundwater flow (by an arrow).

4.1.3 Model setup and calibration

The numerical model MACRO (version 5.1, Larsbo *et al.* (2005)) was applied to the Silstrup site covering the soil profile to a depth of 5 m b.g.s., always including the groundwater table. The model is used to simulate the water flow in the unsaturated zone during the full monitoring period April 2000–June 2008 and to establish an annual water balance.

Compared with the setup in Kjær et al. (2008), a year of validation was added to the MACRO setup for the Silstrup site. The setup was hereby calibrated for the monitoring period May 1999 - June 2004 and validated for the monitoring period July 2004 - June 2008. For this purpose, the following time series have been used: the observed groundwater table measured in the piezometers located in the buffer zone, soil water content measured at three depths (25, 60, and 110 cm b.g.s.) from the two profiles S1 and S2 (Figure 15), and the measured drainage flow. Data acquisition, model setup and results related to simulated bromide transport are described in Barlebo *et al.* (2007) and Rosenbom *et al.* (In prep.). Given impounding of water in the drainage water monitoring well, estimates for the measured drainage on 11 December 2006, 13-14 December, 2006 and 28 February 2007 were based on expert judgment.

4.2 Results and discussion

4.2.1 Soil water dynamics and water balances

The model simulations were largely consistent with the observed data, thus indicating a reasonable model description of the overall soil water dynamics in the unsaturated zone (Figure 16). As in Kjær *et al.* (2008), the simulated groundwater table of this hydraulic year was validated against the much more fluctuating groundwater table measured in piezometer P3, which yielded the best description of measured drainage (Figure 16B and 16C). The earlier reported delay in the initial increase in simulated drainage flow was still present in October and January (Figure 16C). As in the previous monitoring periods, the overall trends in soil water content were described reasonably well (Figure 16D, 16E, and 16F), although the model still tended to describe the subsoil as being much drier during the summer period than measured by the deeper TDR probes (Figure 16E and 16F).

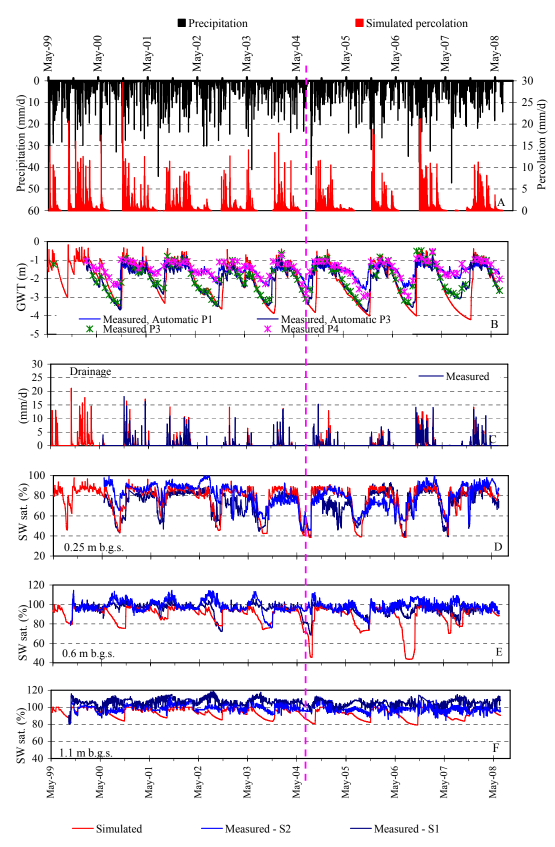


Figure 16. Soil water dynamics at Silstrup: Measured precipitation and simulated percolation 1 m b.g.s. (A), simulated and measured groundwater level GWT (B), simulated and measured drainage flow (C), and simulated and measured soil water saturation (SW sat.) at three different soil depths (D, E and F). The measured data in B derive from piezometers located in the buffer zone. The measured data in D, E and F derive from TDR probes installed at S1 and S2 (see Figure 15). The broken vertical line indicates the beginning of the validation period (July 2004 - June 2008).

Table 8. Annual water balance for Silstrup (mm/year). Precipitation is corrected to the soil surface according to the method of Allerup and Madsen (1979).

	Normal precipitation ²⁾	Precipitation	Actual evapotranspiration	Measured drainage	Simulated drainage	Groundwater recharge ³⁾
1.7.99–30.6.00 ¹⁾	976	1175	457	_	443	275 ⁴⁾
1.7.00-30.6.01	976	909	413	217	232	279
1.7.01-30.6.02	976	1034	470	227	279	338
1.7.02-30.6.03	976	879	537	81	74	261
1.7.03-30.6.04	976	760	517	148	97	94
1.7.04-30.6.05	976	913	491	155	158	267
1.7.05-30.6.06	976	808	506	101	95	201
1.7.06-30.6.07	976	1150	539	361	307	249
1.7.07-30.6.08	976	877	456	200	169	221

¹⁾ The monitoring started in April 2000.

Hydraulic conductivity and water content were measured downstream of the field (close to P1 and P4), but no such data were available upstream, where P3 was located. As mentioned above, the calibration showed that the hydraulic conditions around P3 affected the measurements considerably. In order to model the measured values, the calibrated hydraulic conductivity curves were fitted less well to the measured data from P1 and P4 than from the other sites (Barlebo *et al.*, 2007).

The resulting water balance for Silstrup for the entire monitoring period is shown in Table 8. Compared with the previous eight years, the latest hydraulic year July 2007 - June 2008 was characterised by having the second lowest precipitation, the second lowest simulated actual evapotranspiration and the fourth highest measured drainage. Precipitation in this year was characterized by July, October, January and March being very wet and October and May being very dry (Appendix 4). Due to this precipitation pattern, the simulated percolation pattern of the year July 2007 - June 2008 was represented by continuous percolation throughout the period October - June (Figure 18A), leaving the summer months of 2007 approximately without percolation. The climatic setting of this year gave rise to periods with groundwater table above drainage level causing the fourth largest measured drainage since monitoring started in July 2000 (Figure 18B and 18C). Apart from differences in measured and simulated yearly amount of drainage, the overall pattern of drainage levels in the hydraulic years was captured.

4.2.2 Bromide leaching

The bromide concentrations shown in Figure 17 and Figure 18 relate to the bromide applied in May 2000, as described in previous reports (Kjær *et al.* 2003 & 2004) and further evaluated in Rosenbom *et al.* (In prep) and Barlebo *et al.* (2007).

²⁾ Normal values based on time series for 1961–1990 corrected to soil surface.

³⁾ Groundwater recharge calculated as precipitation - actual evapotranspiration - measured drainage.

⁴⁾ Where drainage flow measurements were lacking, simulated drainage flow was used to calculate groundwater recharge.

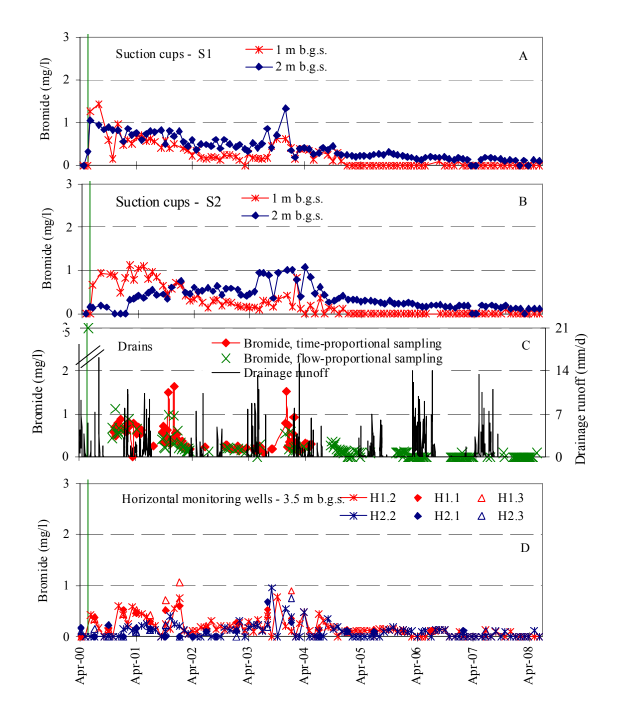


Figure 17. Bromide concentration at Silstrup. A and B refer to suction cups located at S1 and S2. The bromide concentration is also shown for drainage runoff (C) and the horizontal monitoring wells H1 and H2 (D). The green vertical line indicates the date of bromide application.

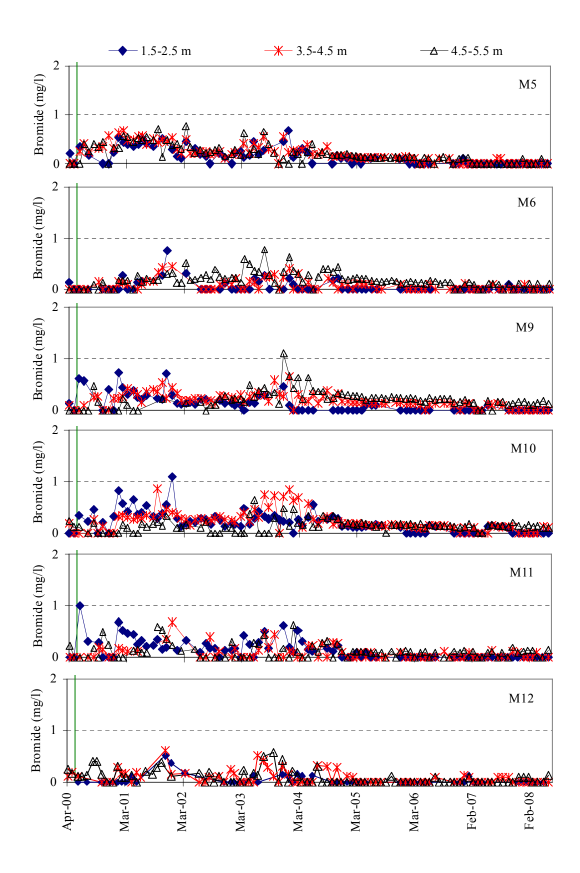


Figure 18. Bromide concentration at Silstrup. The data derive from the vertical monitoring wells (M5–M12). Screen depth is indicated in m b.g.s. The green vertical line indicates the date of bromide application.

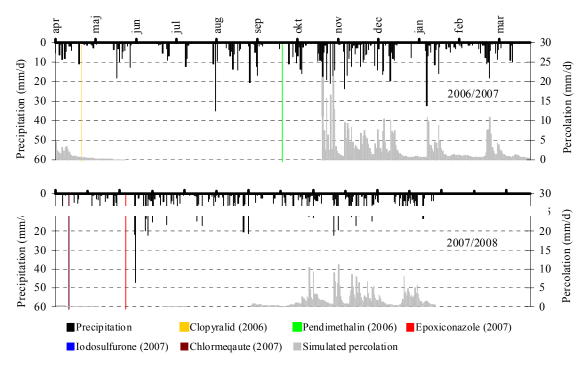


Figure 19. Pesticide application, precipitation and irrigation (primary axis) and simulated percolation 1 m b.g.s. (secondary axis) at Silstrup in 2006/2007 (upper) and 2007/2008 (lower).

4.2.3 Pesticide leaching

Monitoring at Silstrup began in May 2000 and presently encompasses several pesticides and their degradation products, Table 9. Pesticide application during the two most recent growing seasons is shown together with precipitation and simulated precipitation in Figure 19. It should be noted that precipitation in Table 9 is corrected to soil surface according to Allerup and Madsen (1979), whereas percolation (1 m b.g.s.) refers to accumulated percolation as simulated with the MACRO model. It should also be noted that as tribenuronmethyl (applied here as Express) and pyridate (applied here as Lido) degrade rapidly, the leaching risk is more associated with their respective degradation products, triazinamin-methyl and PHCP. For the same reasons it is the degradation products and not the parent compounds that are monitored in the PLAP (Table 9).

Table 9. Pesticides analysed at Silstrup with the product used shown in parentheses. Degradation products are in italics. Precipitation (prec.) and percolation (perc.) are accumulated from date of first application until end of monitoring. 1^{st} month perc. refers to accumulated percolation within the first month after application. C_{mean} refers to average leachate concentration in the drainage water within the first drainage season after application (See Appendix 2 for calculation methods). The number of pesticide-positive samples is indicated in parentheses.

Crop and analysed pesticides	Application	End of	Prec.	Perc.	1 st month	C_{mean}
	date	monitoring	(mm)	(mm)	perc. (mm)	$(\mu g/l)$
Fodder beet 2000						
Metamitron (Goltix WG)	May 00	Apr 03	2634	1328	53	0.05 (69)
- metamitron-desamino	,	1				0.06 (61)
Ethofumesate (Betanal Optima)	May 00	Apr 03	2634	1328	53	0.03 (24)
Desmedipham (Betanal Optima)	May 00	Apr 03	2634	1328	53	< 0.01(1)
- EHPC	,	•				< 0.02 (0)
Phenmedipham (Betanal Optima)	May 00	Apr 03	2634	1328	53	< 0.01 (0)
- MHPC	-	-				< 0.02 (0)
-3-aminophenol						< 0.02 (0)
Fluazifop-P-butyl (Fusilade X-tra)	Jun 00	Jul 02	1953	1019	5	< 0.01 (0)
- fluazifop (free acid)						< 0.02(1)
Pirimicarb (Pirimor G)	Jul 00	Jul 07	6452	2825	1	< 0.01 (17*)
- pirimicarb-desmethyl						<0.01 (1*)
- pirimicarb-desmethyl-formamido						< 0.02 (0)
Spring barley 2001						
Triazinamin-methyl 1) (Express)	May 01	Jul 03	1941	951	10	< 0.02 (0)
Flamprop-M-isopropyl (Barnon Plus 3)	Jun 01	Jul 03	1928	944	3	< 0.01 (13)
- flamprop (free acid)						< 0.01 (7)
Propiconazole (Tilt Top)	Jun 01	Jul 03	1928	944	3	< 0.01 (6)
Fenpropimorph (Tilt Top)	Jun 01	Jul 03	1928	944	3	< 0.01 (0)
- fenpropimorphic acid						< 0.01 (1)
Dimethoate (Perfekthion 500 S)	Jul 01	Jul 03	1882	937	3	0.02(2)
Maize 2002						
Glyphosate (Roundup Bio)	Oct 01	Apr 06	3802	1694	44	<0.13 (71*)
- AMPA		1				$0.06(137^*)$
<i>PHCP</i> ²⁾ (Lido 410 SC)	May 02	Jul 04	1764	738	6	0.06 (18)
Terbuthylazine (Lido 410 SC)	May 02	Apr 06	3320	1327	6	0.07 (96)
- desethyl-terbuthylazine	,	Apr 05				0.15 (269)
- 2- hydroxy-terbuthylazine		Apr 05				$^{3)}(26)$
- 2-hydroxy-desethyl-terbuthylazine		Apr 05				³⁾ (29)
- desisopropyl-atrazine		Apr 05				³⁾ (47)

¹⁾ Degradation product of tribenuron-methyl. The parent compound degrades too rapidly to be detected by monitoring.

The leaching risk of pesticides applied during 2000, 2001, 2002, 2003, 2004 and 2005 has been evaluated in Kjær *et al.* (2002, 2003, 2004, 2005c, 2007 & 2008), respectively.

The herbicide propyzamide applied late autumn/beginning of the winter 2005, together with its three metabolites RH24580, RH 24644 and RH24655 has now been monitored for nearly 2.5 years,—The leaching from the root zone to the drainage system was in particular related to propyzamide, followed by RH24644 and RH 24580 (see Figure 23 in Kjær *et al.* (2008)). The metabolite RH24655 was not detected at all. Only propyzamide concentrations in drainage water exceeded the 0.1μg/l limit. One metabolite, RH24644, was detected in groundwater on 1 December 2005 at 0.032 μg/l in H1 (3.5 m b.g.s.) and 0.016 in M5 (1.5-2.5 m b.g.s.) (data not shown) Propyzamide was found in groundwater nine times. The limit of 0.1μg/l was exceeded twice, both samples taken on 1 December 2005 (0.14 μg/l in H1 (3.5 m b.g.s.) and 0.11μg/l in M5 (1.5-2.5 m b.g.s.)). Further information and illustration of these monitoring results can be found in Kjær et al. (2008).

²⁾ Degradation product of pyridate. The parent compound degrades too rapidly to be detected by monitoring.

³⁾ Average leachate concentration within the first drainage season after application could not be calculated, as monitoring started January 2003 (7 mount after application). See Kjær et al.(2007) for further information

^{*} Pesticide has been applied more than once, and the findings are not necessarily related to one specific application.

Table 9 continued. Pesticides analysed at Silstrup with the product used shown in parentheses. Degradation products are in italics. Precipitation (prec.) and percolation (perc.) are accumulated from date of first application until end of monitoring. 1^{st} month perc. refers to accumulated percolation within the first month after application. C_{mean} refers to average leachate concentration in the drainage water within the first drainage season after application (See Appendix 2 for calculation methods). The number of pesticide-positive samples is indicated in parentheses.

Crop and analysed pesticides	Application	End of	Prec.	Perc.	1 st month	C _{mean}
	date	monitoring	(mm)	(mm)	Perc. (mm)	(μg/l)
Peas 2003						
Bentazone (Basagran 480)	May 03	Jul 06	2634	1055	44	0.26 (52)
- AIBA						< 0.01 (0)
Pendimethalin (Stomp SC)	May 03	Apr 06	2634	1055	44	< 0.01 (0)
Glyphosate (Roundup Bio)	Sep 03	Apr 06	2207	971	0	<0.01 (71*)
- AMPA						$0.02(137^*)$
Winter wheat 2004						
Prosulfocarb (Boxer EC)	Oct 03	Apr 06	2125	974	37	0.01(6)
MCPA (Metaxon)	May 04	Jul 06	1797	710	4	< 0.01 (0)
- 4-chlor,2-methylphenol						<0.01 (0)
Azoxystrobin (Amistar)	Jun 04	Jul 06	1781	706	0	0.01 (6*)
- CyPM		Jul 07	2931	1202	0	0.09 (62*)
Pirimicarb (Pirimor G)	Jul 04	Jul 07	2818	1205	0	<0.01 (17*)
- pirimicarb-desmethyl						<0.01 (1*)
- pirimicarb-desmethyl-formamido						<0.02(0)
Spring barley 2005						
Fluroxypyr (Starane 180 S)	May 05	Jul 07	2012	830	11	<0.02 (0)
Azoxystrobin (Amistar)	Jun 05	Jul 06	862	332	10	0.01 (6*)
- CyPM	Jun 05	Jul 07	2012	828	10	0.02 (62*)
Pirimicarb (Pirimor G)	Jul 05	Jul 07	1933	818	0	<0.01 (17*)
- pirimicarb-desmethyl						<0.01 (1*)
- pirimicarb-desmethyl-formamido						<0.01 (0*)
Winter rape 2006						7)
Propyzamide (Kerb 500 SC)	Nov 05	Apr 08	2345	1115	75	$0.22(32)^{1}$
- RH-24644						$0.01(17)^{1}$
- RH-24580						<0.01 (2) ¹⁾
- RH-24655			2000	0.50	0	<0.01 (0) 1)
Clopyralid (Matrigon)	Apr 06	Apr 08	2009	859	8	<0.01 (0)
Winter wheat 2007						
Pendimethalin (Stomp Pentagon)	Sep 06	Apr 08	1686	865	0	<0.04 (14)
Iodosulfuron-methyl-sodium (Hussar	Apr 07	Jul 08 [†]	1099	392	3	<0.01 (0)
OD)						<0.01 (0)
- metsulfuron-methyl						<0.01 (0)
-triazinamin						
Chlormequat (Cycocel 750)	Apr 07	Jul 08 [†]	1099	392	3	<0.01(1)
Epoxiconazole (Opus)	Jun 07	Jul 08 [†]	1026	390	0	< 0.01 (0)

^{*} Pesticide has been applied more than once, and the findings are not necessarily related to one specific application.

Clopyralid, applied on 20 April 2006, was not found in any of the water samples taken within a two-year period (Table 9).

The herbicide pendimethalin, applied on 22 September 2006, was found in drainage water 14 times, starting two months after the application and the last time 4.5 months after the application, Figure 20, in concentrations not exceeding 0.1 μ g/l. Pendimethalin was not detected in groundwater (Table A5.3 in Appendix 5).

Apart from one detection of chlormequat (drainage water samples collected 14 February 2007), none of the pesticides used on winter wheat in 2006 have yet been found in any of the analysed water samples. The final evaluation awaits an additional year of monitoring.

¹⁾ Drainage runoff commenced two weeks prior to the application of propyzamide, and the weighted concentrations refer to the period from the date of application until 1 July 2007.

[†] Monitoring will continue for an additional year. The values for prec. and perc. are accumulated up to July 2007.

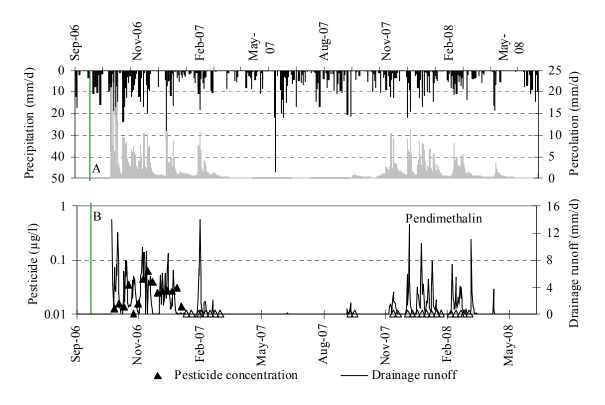


Figure 20. Precipitation and simulated percolation 1 m b.g.s. (A) together with the concentration of pendimethalin (B) in the drainage runoff at Silstrup. The green vertical line indicates the date of pendimethalin application. Open diamonds indicate values below the detection limit of $0.01 \mu g/l$.

5 Pesticide leaching at Estrup

5.1 Material and methods

5.1.1 Site description and monitoring design

Estrup is located in central Jutland (Figure 1) west of the Main Stationary Line on a hillisland, i.e. a glacial moraine preserved from the Weichselian Glaciation. Estrup has thus been exposed to weathering, erosion, leaching and other geomorphological processes for a much longer period than the other sites. The test field covers a cultivated area of 1.26 ha (105 x 120 m) and is virtually flat (Figure 25). The site is highly heterogeneous with considerable variation in both topsoil and aquifer characteristics (Lindhardt et al. (2001). Such heterogeneity is quite common for this geological formation, however. Based on three profiles excavated in the buffer zone bordering the field the soil was classified as Abruptic Argiudoll, Aquic Argiudoll and Fragiaquic Glossudalf (Soil Survey Staff, 1999). The topsoil is characterized as sandy loam with a clay content of 10–20% and an organic carbon content of 1.7–7.3%. A C-horizon of low permeability also characterizes the site. The saturated hydraulic conductivity in the C-horizon is 10⁻⁸ m/s, which is about two orders of magnitude lower than that at the other loamy sites (Table 1). The geological structure is complex comprising a clay till core with deposits of different age and composition (Lindhardt et al. (2001). A brief description of the sampling procedure is provided in Appendix 2. The monitoring design and test site are described in detail in Lindhardt et al. (2001) and the analysis methods in Kjær et al. (2002). Please note that the geological conditions only allowed one of the planned horizontal wells to be installed as drilling in sand proved impossible.

5.1.2 Agricultural management

Management practice during the 2005 - 2007 growing seasons is briefly summarized below and detailed in Appendix 3 (Table A3.4). For information about management practice during the previous monitoring periods see Kjær *et al.* (2002, 2003, 2004, 2005c, 2007 & 2008).

Ploughing of the field took place on 12 April 2006 and on 27 April spring barley was sown (cv. Simba), which emerged on 6 May. Weeds were treated twice with fluroxypyr, 17 May and 2 June, but the substance was not included in the monitoring. On 6 June, at the beginning of stem elongation, the herbicide florasulam was applied. On 29 June at 70% emergence of inflorescence, the fungicide azoxystrobin was used. The amount of grain harvest on 17 August was 59.2 hkg/ha, a roughly average yield (Plantedirektoratet, 2006). Straw, 26.8 hkg/ha, was removed on 11 September and two days later the field was ploughed. Winter wheat (cv. Smuggler) was sown on 14 September and emerged on 22 September 2007. When on 11 October the crop had three leaves unfurled, the herbicides mesosulfuron and iodosulfuron (Atlantis WG) were applied. To prevent lodging, plant growth was regulated using chlormequat on 11 April 2007 at the beginning of stem elongation. The fungicide epoxiconazole was used on 31 May when 70% of the inflorescence had emerged. On 7 August the amount of 81, 5

hkg/ha of grain (85% dry matter) was harvested, and 47.7 hkg/ha of straw (100% dry matter) was shredded the subsequent day. Yields of grain were similar for this soil type and year (Plantedirektoratet, 2008).

The field was ploughed on 2 October 2007 and the following day sown with winter wheat (cv. Frument). Due to the late sowing, the crop did not emerge until 17 October. When on 30 October the crop had two leaves unfurled, it was sprayed with the herbicide picolinafen, and on 22 November with the fungicide tebuconazole when it had three leaves unfurled.

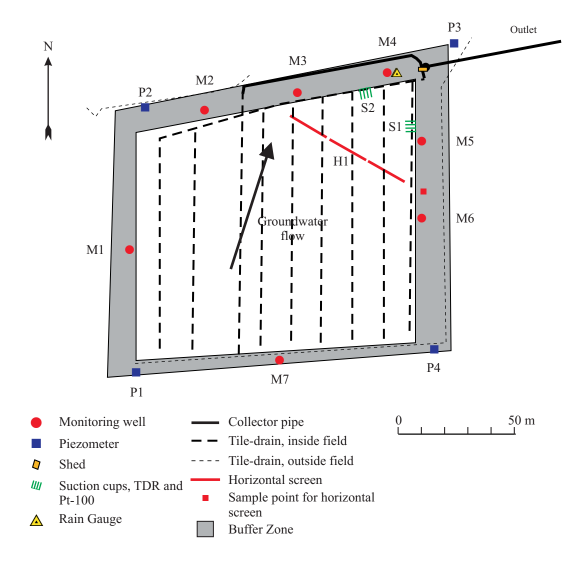


Figure 21. Overview of the Estrup site. The innermost white area indicates the cultivated land, while the grey area indicates the surrounding buffer zone. The positions of the various installations are indicated, as is the direction of groundwater flow (by an arrow).

5.1.3 Model setup and calibration

The numerical model MACRO (version 5.1, Larsbo *et al.* (2005)) was applied to the Estrup site covering the soil profile to a depth of 5 m b.g.s., always including the groundwater table. The model is used to simulate the water flow in the unsaturated zone during the monitoring period from July 2000 - June 2008 and to establish an annual water balance.

Compared to the setup in Kjær *et al.* (2008), a year of validation was added to the MACRO setup for the Estrup site. The setup was hereby calibrated for the monitoring period May 1999 - June 2004 and validated for the monitoring period July 2004 - June 2008. For this purpose, the following time series have been used: the observed groundwater table measured in the piezometers located in the buffer zone, measured drainage flow and soil water content measured at two depths (25 and 40 cm b.g.s.) from the soil profile S1 (Figure 25). The TDR probes installed at the other depths yielded unreliable data with saturations far exceeding 100% and dynamics with increasing soil water content during the drier summer periods (data not shown). No explanation can presently be given for the unreliable data, and they have been excluded from the analysis. The data from the soil profile S2 have also been excluded due to a problem of water ponding above the TDR probes installed at S2, as mentioned in Kjær *et al.* (2003). Because of the erratic TDR data, calibration data are limited at this site. Data acquisition, model setup as well as results related to simulated bromide transport are described in Barlebo *et al.* (2007) and Rosenbom *et al.* (in prep.).

5.2 Results and discussion

5.2.1 Soil water dynamics and water balances

The model simulations were generally consistent with the observed data (which were limited compared to other PLAP sites, as noted above), indicating a good model description of the overall soil water dynamics in the unsaturated zone (Figure 22). The model provided an acceptable simulation of the overall level of the groundwater table. A drop in the measured groundwater table was seen after short periods of low precipitation (Figure 22B). The simulated groundwater table did not seem as sensitive to these short periods of low precipitation and tended not to drop as much as the measured values. Since the TDR data are limited in the subsoil, a more detailed study of dynamics in these layers is difficult. However, the overall soil water saturation at 25 and 40 cm b.g.s. was captured excluding periods with drastic drops in the level of saturation (Figure 26D and 26E). Especially the size of the water saturation drop in May-June 2008 was not captured. Nothing special is noted for the groundwater table in the latest monitoring period (July 2007 - June 2008). As in previous years (Kjær *et al.*, 2008), the simulated groundwater table often fluctuates slightly above the drain depth during periods of drainage flow.

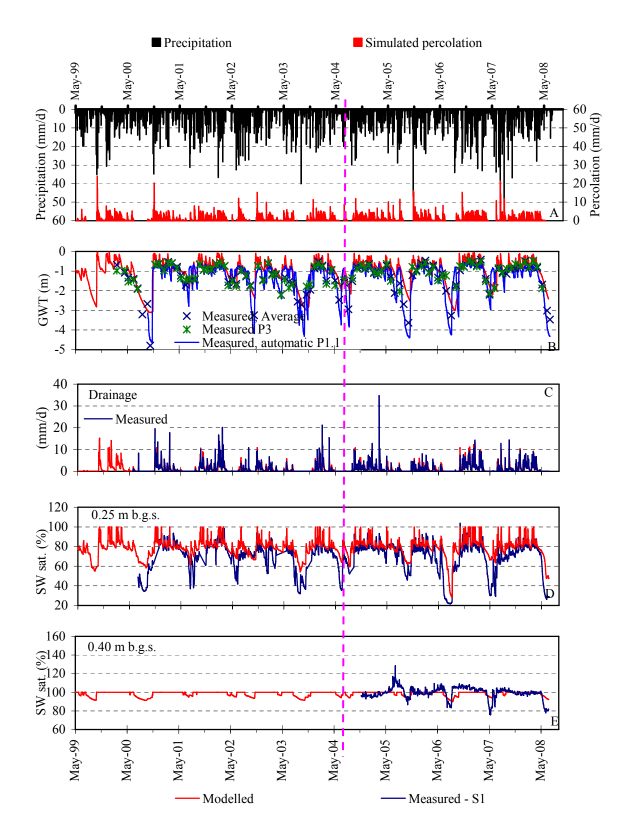


Figure 22. Soil water dynamics at Estrup: Measured precipitation and simulated percolation 0.6 m b.g.s. (A), simulated and measured groundwater level (B), simulated and measured drainage flow (C), and simulated and measured soil saturation (SW sat.) at two different soil depths (D and E). The measured data in B derive from piezometers located in the buffer zone. The measured data in D and E derive from TDR probes installed at S1 (see Figure 21). The broken vertical line indicates the beginning of the validation period (July 2004 - June 2008).

Table 10. Annual water balance for Estrup (mm/year). Precipitation is corrected to the soil surface according to the method of Allerup and Madsen (1979).

	Normal precipitation ²⁾	Precipitation	Actual evapotranspiration	Measured drainage	Simulated drainage	Groundwater recharge ³⁾
1.7.99–30.6.00 ¹⁾	968	1173	466	_	553	154 ⁴⁾
1.7.00-30.6.01	968	887	420	356	340	111
1.7.01-30.6.02	968	1290	516	505	555	270
1.7.02-30.6.03	968	939	466	329	346	144
1.7.03-30.6.04	968	928	499	298	312	131
1.7.04-30.6.05	968	1087	476	525	468	86
1.7.05-30.6.06	968	897	441	258	341	199
1.7.06-30.6.07	968	1365	515	547	618	303
1.7.07-30.6.08	968	1045	478	521	556	46

¹⁾ Monitoring started in April 2000.

The simulated drainage (Figure 22C) matched the measured drainage flow quite well. Drainage runoff over the whole monitoring period was high compared to that of the other two till sites being investigated in the PLAP. This was due to a significantly lower permeability of the C-horizon than of the overlying A and B horizons (see Kjær *et al.* 2005c for details).

The resulting water balance for Estrup for the nine monitoring periods is shown in Table 10. Compared with the previous eight years, the latest hydraulic year July 2007 - June 2008 was characterised by having the 5th highest precipitation, the 3rd highest simulated actual evapotranspiration and the 2nd highest measured and simulated drainage. Precipitation in this year was characterized by July and Marts being very wet and October and April-June being very dry (Appendix 4). Due to this precipitation pattern, the simulated percolation pattern of the year July 2007 - June 2008 was similar to the pattern of earlier years in the monitoring period, and was represented by continuous percolation throughout the period September-May (Figure 26A), leaving the summer months with scattered periods of both percolation and drainage runoff (Figure 26B and 26C). Apart from differences in measured and simulated yearly amount of drainage, the overall trend in drainage amount pattern of the hydraulic years was captured.

5.2.2 Bromide leaching

Bromide has now been applied twice at Estrup. The bromide concentrations measured up to October 2005 (Figure 23 and Figure 24) relate to the bromide applied in spring 2000, as described further in Kjær *et al.* (2003). Leaching of the bromide applied in November 2005 is evaluated in Rosenbom *et al.* (In prep.).

²⁾ Normal values based on time series for 1961–1990 corrected to the soil surface.

³⁾ Groundwater recharge is calculated as precipitation minus actual evapotranspiration minus measured drainage.

⁴⁾ Where drainage flow measurements are lacking, simulated drainage flow was used to calculate groundwater recharge.

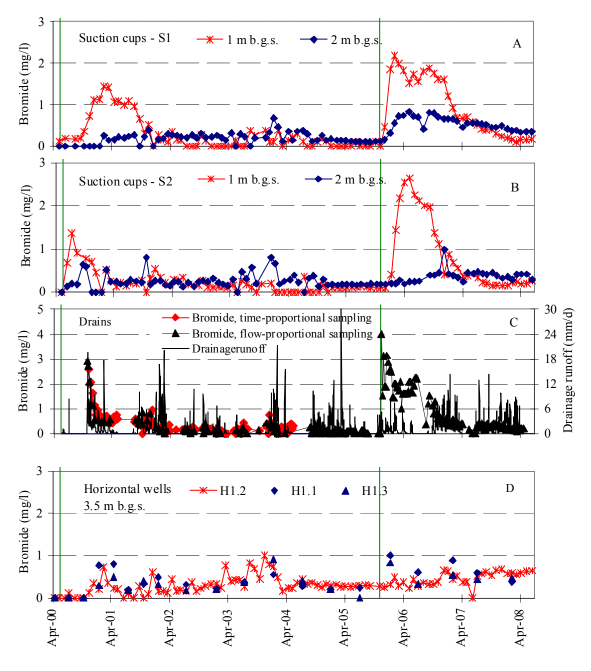


Figure 23. Bromide concentration at Estrup. A and B refer to suction cups located at S1 and S2, respectively. The bromide concentration is also shown for drainage runoff (C) and the horizontal monitoring well H1 (D). The green vertical lines indicate the dates of bromide applications.

5.2.3 Pesticide leaching

Monitoring at Estrup began in May 2000 and encompass several pesticides and their degradation products, as indicated in Table 11. Pesticide application during the two most recent growing seasons (2006-2007) is shown together with precipitation and simulated precipitation in Figure 25. It should be noted that precipitation is corrected to the soil surface according to Allerup and Madsen (1979), whereas percolation (0.6 m b.g.s.) refers to accumulated percolation as simulated with the MACRO model (Section 5.2.1). Moreover, pesticides applied later than April 2008 are not evaluated in this report and hence are not included in Table 11.

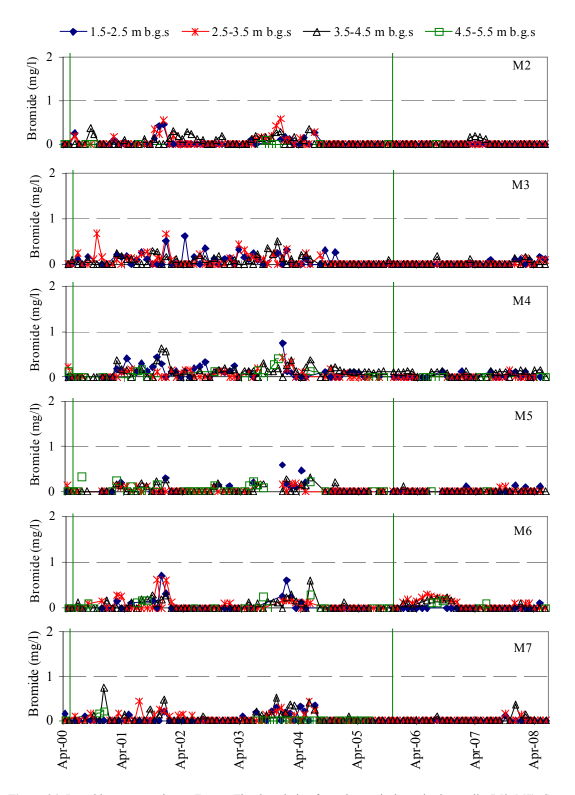


Figure 24. Bromide concentration at Estrup. The data derive from the vertical monitoring wells (M2–M7). Screen depth is indicated in m b.g.s. The green vertical lines indicate the dates of bromide applications.

Table 11. Pesticides analysed at Estrup with the product used shown in parentheses. Degradation products are in italics. Precipitation (prec.) and percolation (perc.) are accumulated from the date of first application until the end of monitoring. 1st month perc. refers to accumulated percolation within the first month after application. C_{mean} refers to average leachate concentration in the drainage water within the first drainage season after application (See Appendix 2 for calculation methods). The number of pesticide-positive samples is indicated in parentheses.

Crop and analysed pesticides	Application	End of	Prec.	Perc.	1st month	C _{mean}
	date	monitoring	(mm)	(mm)	Perc. (mm)	$(\mu g/l)$
Spring barley 2000						
Metsulfuron-methyl (Ally)	May 00	Apr 03	2990	1456	29	< 0.01(1)
- triazinamin	,	•				< 0.02(1)
Flamprop-M-isopropyl (Barnon Plus 3)	May 00	Apr 03	2914	1434	2	0.02(20)
- flamprop (free acid)	-	•				0.01(13)
Propiconazole (Tilt Top)	Jun 00	Apr 05	4938	2294	0	$0.01(27^*)$
Fenpropimorph (Tilt Top)	Jun 00	Jul 02	2211	1048	0	< 0.01(1)
- fenpropimorphic acid						< 0.02(0)
Dimethoate (Perfekthion 500 S)	Jun 00	Jul 02	2211	1048	0	< 0.01(0)
Pea 2001						
Glyphosate (Roundup Bio)	Oct 00	Jul 08 [†]	8226	4037	123	$0.54(260^*)$
- AMPA						0.17(316*)
Bentazone (Basagran 480)	May 01	Jul 08	7629	3621	9	0.03 (139)
- AIBA						<0.01 (2)
Pendimethalin (Stomp SC)	May 01	Jul 03	2208	1096	9	< 0.01(37*)
Pirimicarb (Pirimor G)	Jun 01	Jul 05	4251	1995	10	$0.01(40^*)$
- pirimicarb-desmethyl						< 0.02(0)
- pirimicarb-desmethyl-formamido						< 0.02(26*)
Winter wheat 2002						
Ioxynil (Oxitril CM)	Nov 01	Jul 03	1580	860	52	$0.04(20)^1$
Bromoxynil (Oxitril CM)	Nov 01	Jul 03	1580	860	52	$0.01(3)^{1}$
Amidosulfuron (Gratil 75 WG)	Apr 02	Jul 04	2148	928	8	<0.01(0)
MCPA (Metaxon)	May 02	Jul 04	2091	928	0	< 0.01(12)
- 4-chlor,2-methylphenol	-					< 0.01(1)
Propiconazole (Tilt 250 EC)	May 02	Apr 05	2920	1336	39	$0.02(27^*)$
Pirimicarb (Pirimor G)	Jun 02	Jul 05	2982	1403	58	$0.01(40^*)$
- pirimicarb-desmethyl						< 0.02(0)
- pirimicarb-desmethyl-formamido		Apr 06				< 0.02(26*)

Systematic chemical nomenclature for the analysed pesticides is given in Appendix 1.

The leaching risk of pesticides applied during 2000, 2001, 2002, 2003, 2004 and 2005 has been evaluated in Kjær et al. (2002, 2003, 2004, 2005c, 2007 & 2008).

Since metabolites of terbuthylazine (applied in 2005) and glyphosate as well as its metabolite AMPA (applied 2005) were still included in the current monitoring period, the results of these applications are summarised below. For a detailed description of the leaching pattern, including primary data and climate condition characterising the monitoring periods, reference is made to Kjær et al. (2007).

^{*}Pesticides have been applied twice, and the findings are not necessarily related to one specific application.

[†] Monitoring will continue during the following year. The values for prec. and perc. are accumulated up to July 2006.

1) Drainage runoff commenced about two and a half months prior to the application of ioxynil and bromoxynil, and the weighted concentrations refer to the period from the date of application until 1 July 2002.

Table 11 continued. Pesticides analysed at Estrup with the product used shown in parentheses. Degradation products are in italics. Precipitation (prec.) and percolation (perc.) are accumulated from the date of first application until the end of monitoring. 1st month perc. refers to accumulated percolation within the first month after application. C_{mean} refers to average leachate concentration in the drainage water within the first drainage season after application (See Appendix 2 for calculation methods). The number of pesticide-positive samples is indicated in parentheses.

Crop and analysed pesticides	Application	End of	Prec.	Perc.	1 st month	C _{mean}
Crop and analysed pesticides	date	monitoring	(mm)	(mm)	perc. (mm)	(μg/l)
Fodder beet 2003					•	
Glyphosate (Roundup Bio)	Sep 02	Jul 08 [†]	6032	2961	0	$0.43~(260^*)$
- AMPA	•					$0.19(316^*)$
Ethofumesate (Betanal Optima)	May 03	Apr 06	2901	1371	50	0.11(35)
Metamitron (Goltix WG)	May 03	Apr 06	2901	1371	50	1.1(42)
- metamitron-desamino						0.21(49)
Pirimicarb (Pirimor G)	Jul 03	Jul 05	2071	939	0	<0.01(40*)
- pirimicarb-desmethyl		Jul 05				<0.01(0)
 pirimicarb-desmethyl-formamido 		Apr 06				$0.12(26^*)$
Spring barley 2004						
Fluroxypyr (Starane 180)	May 04	Jul 06	2073	1030	0	< 0.02 (2)
Azoxystrobin (Amistar)	Jun 04	Jul 08	4452	2209	38	0.12 (65*)
- CyPM						0.23 (132*)
Maize 2005						
Terbuthylazine (Inter-Terbuthylazin)	May 05	Jul 08 [†]	3338	1628	32	0.48 (100)
- desethyl-terbuthylazine		Jul 08 [†]				0.31 (127)
- 2-hydroxy-terbuthylazine		Jul 08 [†]				0.11 (87)
- desisopropyl-atrazine		Jul 08				0.02 (97)
 2-hydroxy-desethyl-terbuthylazine 		Jul 08				0.24 (86)
Bentazone (Laddok TE)	Jun 05	Jul 08	3338	1628	10	0.18 (139)
- AIBA		±				<0.01 (2)
Glyphosate (Roundup Bio)	Nov 05	Jul 08 [†]	2993	1521	68	$4.04 (260^*)^{1}$
- AMPA						$0.42 (316^*)^{I}$
Spring barley 2006						
Florasulam (Primus)	Jun 06	Jul 08	2442	1163	0	< 0.01 (0)
- florasulam-desmethyl						< 0.03 (0)
Azoxystrobin (Amistar)	Jun 06	Jul 08	2414	1170	0	$0.03~(65^*)$
- CyPM						$0.13 (132^*)$
Winter wheat 2007						
Mesosulfuron-methyl (Atlantis WG)	Oct 06	Jul 08	2059	1095	63	<0.01 (13) ¹⁾
- mesosulfuron						< 0.02(0)
Chlormequat (Cycocel 750)	Apr 07	Jul 08 [†]	1337	603	0	< 0.01 (1)
Epoxiconazole (Opus)	May 07	Jul 08	1199	600	45	0.01 (13)
Winter wheat 2008		±				•.
Picolinafen (Pico 750 WG)	Oct 07	Jul 08 [†]	613	381	52	$0.03(17)^{1}$
- CL153815						$0.24 (21)^{1}$
Tebuconazol (Folicur EC 250) Systematic chemical nomenclature for the	Nov 07	Jul 08 [†]	565	345	77	0.43 (23) 1)

Systematic chemical nomenclature for the analysed pesticides is given in Appendix 1.

Terbuthylazine was applied during the growing of maize in 2005. Terbuthylazine and all of its measured metabolites leached to the drainage system (Figure 26). In the third year after application they were all still found in drainage water samples, but concentrations were in nearly all instances less than 0.1 µg/l. Concentrations in drainage water on average exceeded 0.1 µg/l for all but desisopropyl-atrazine (Table 11). In addition to being found in drainage water, the metabolite desisopropyl-atrazine was frequently detected in groundwater, but in low concentrations (Figure 27B, Table A5.4 in Appendix 5). Another metabolite, desethyl-terbuthylazine, was found five times in low concentrations in groundwater (Figure 27C and table A5.4 in Appendix 5).

^{*} Pesticide has been applied more than once, and the findings are not necessarily related to one specific application.

[†] Monitoring will continue during the following year. The values for prec. and perc. are accumulated up to July 2006.

Drainage runoff commenced prior to the application of pesticide and the weighted concentrations refer to the period from the date of application until 1 July the following year.

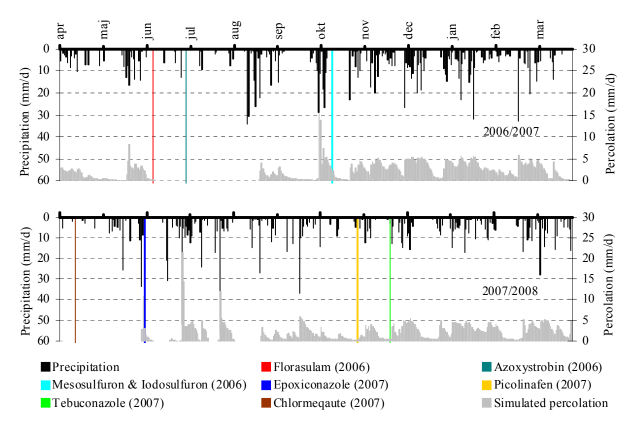


Figure 25. Pesticide application and precipitation (primary axis) together with simulated percolation 0.6 m b.g.s. (secondary axis) at Estrup in 2006/2007 (upper) and 2007/2008 (lower).

The herbicide glyphosate has now been applied at Estrup in 2000, 2002 and 2005 (Figure 28). The most recent application (November 2005) was followed by a significantly higher leaching of glyphosate into the drainage system compared to the two previous applications. This time a concentration peak of $31.0 \,\mu\text{g/l}$ was found eight days after the spraying, and a concentration above $0.1 \,\mu\text{g/l}$ was upheld for a period of six months (Figure 28 B). The average concentration of glyphosate following the 2005 application was $4.04 \,\mu\text{g/l}$, which is 7-9 times higher than following the 2000 and 2002 applications (Table 11). The average concentration of AMPA in the drainage water following the 2005 application amounted to $0.42 \,\mu\text{g/l}$ (Table 11), which was $2.2-2.5 \,\mu\text{times}$ higher than after the previous two applications (Table 11).

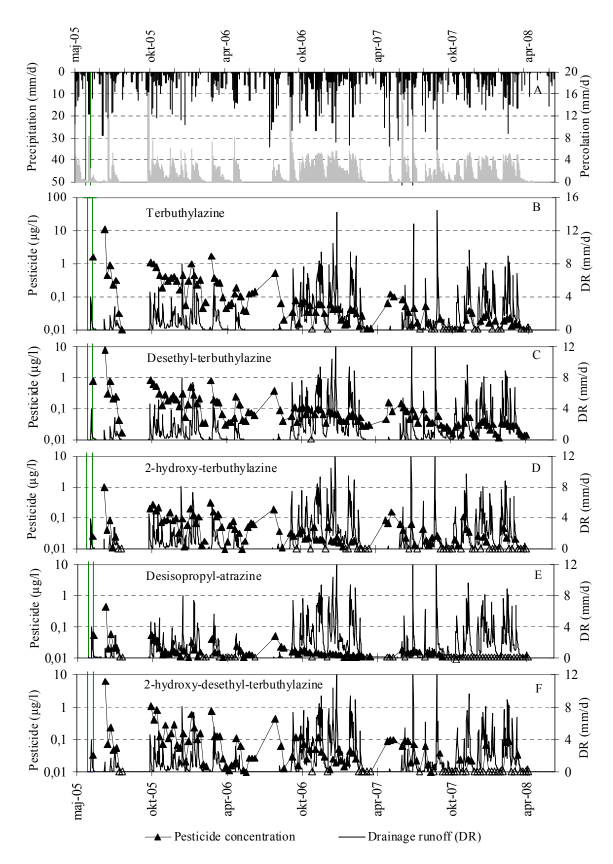


Figure 26. Precipitation and simulated percolation (A) together with concentration of terbuthylazine (B), desethylterbuthylazine (C), 2-hydroxyl-terbuthylazine (D), desisopropyl-atrazine (E) and 2-hydroxy-desethyl-terbuthylazine (F) in the drainage runoff (DR on the secondary axis) at Estrup. The green vertical lines indicate the dates of applications. Open triangles indicate values below the detection limit of $0.01~\mu g/l$.

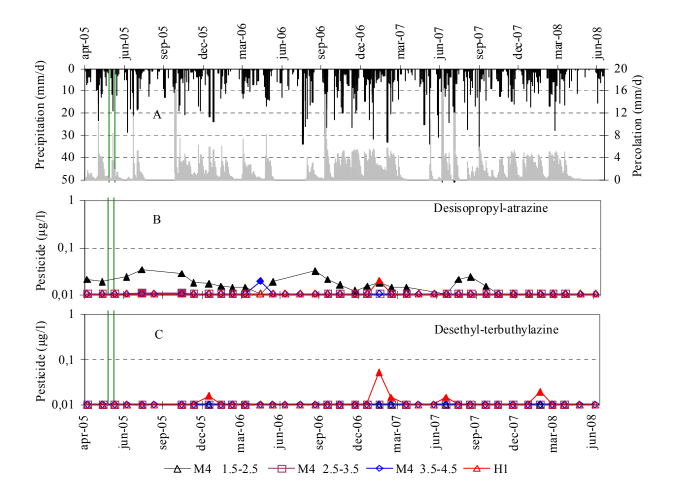


Figure 27. Precipitation and simulated percolation (A) together with concentration of desisopropyl-atrazine (B) and desethyl-terbuthylazine (C) in groundwater monitoring screens at Estrup in 2005/2008. The green vertical lines indicate the dates of applications, and numbers in parentheses the screen depth.

Florasulam, applied on 6 June 2006, and its metabolite florasulam-desmethyl has not been detected in any of the water samples analyzed. Azoxystrobin, applied on 9 June 2006, and its metabolite CyPM leached to the depth of the drainage system in average concentrations of 0.03 µg/l for azoxystrobin and 0.13 µg/l for CyPM (Table 11). A maximum concentration of 1.4 µg/l azoxystrobin was found 2.5 months after application (Figure 29B). This finding coincided with some heavy rain events (Figure 29A). The maximum measured concentration of CyPM of 0.66 µg/l detected on 24 August was less than that of azoxystrobin, but CyPM seemed to be more persistent (Figure 29C). While leaching of azoxystrobin ceased one year after application (the last detection was on 23 August 2007), leaching of the metabolite CyPM continued. Elevated concentrations of CyPM were seen toward the end of this monitoring period (e.g. two years after application), which indicates that leaching of CyPM has not yet ceased. Apart from two detections of CyPM, neither CyPM nor azoxystrobin have been detected in samples collected from the groundwater monitoring screen (Table A5.4 in Appendix 5).

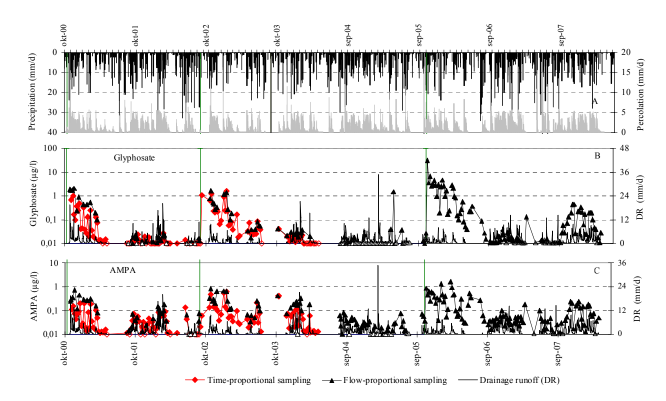


Figure 28. Precipitation and simulated percolation (A) together with the concentration of glyphosate (B) and AMPA (C) in the drainage runoff (DR. on the secondary axis) at Estrup. Data represent a seven-year period including three applications of glyphosate as indicated by the green vertical lines. Open diamonds and triangles indicate values below the detection limit of $0.01~\mu g/l$.

The leaching risk of pesticides applied in autumn 2007 and 2008 will not be evaluated until the 2009 monitoring results become available, i.e. when two years of monitoring data have been collected. However, the following pesticides leached to drainage water, whereas there has been no leaching to groundwater screens situated beneath the drainage system:

- Epoxiconazole, applied on 31 May 2007 (Figure 30B). Concentrations were in two instances above 0.1 μ g/l (Figure 30B), but the average first-year concentration was 0.01 μ g/l (Table 11). The most recent detection of epoxiconazole was on 13 March 2008.
- Picolinafen was applied on 30 October 2007. Concentrations in drainage water never exceeded 0.1 μ g/l (Figure 30C). However the metabolite of picolinafen, CL153815, exceeded the 0.1 μ g/l in several instances (Figure 30D).
- Tebuconazole, applied on 22 October 2007, was seen in drainage water leachate on several occasions, reaching a maximum of 1.5 μ g/l on 29 November 2007, and most drainage water samples analysed contained concentrations of tebuconazole above 0.1 μ g/l (Figure 30E).

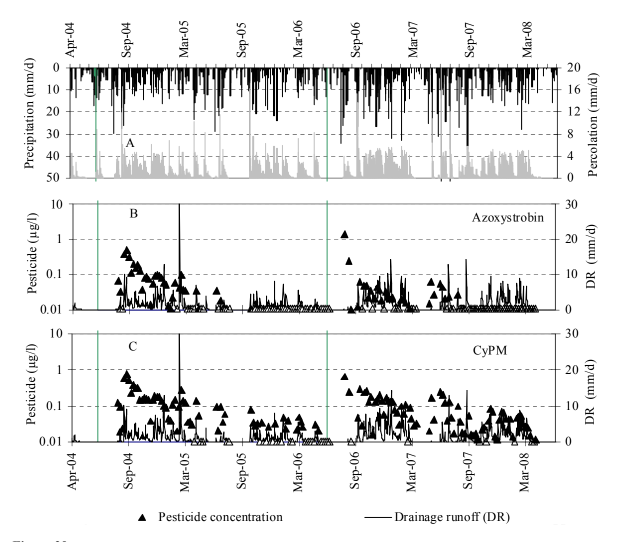


Figure 29. Precipitation and simulated percolation (A) together with concentration of azoxystrobin (B) and CyPM (C) in the drainage runoff (DR on the secondary axis) at Estrup in 2004/2008. The green vertical lines indicate the date of application. Open diamonds and triangles indicate values below the detection limit of $0.01 \,\mu g/l$.

Pesticide leaching at Estrup is generally confined to the depth of the drainage system. Apart from bentazone, glyphosate and desisopropylatrazin being detected in 12, 25 and 26 groundwater samples, pesticides have only sporadically been detected in groundwater monitoring screens below the depth of the drainage system (Appendix 5, Table A5.4). Due to decreased hydraulic conductivity and a lower degree of preferential flow, transport of water and solutes at Estrup are much slower beneath the drainage system than above it. Slow transport may allow for dispersion, dilution, sorption and degradation, thereby further reducing the deep transport. Compared to the other loamy soils investigated, the retention characteristics at Estrup suggest that the C-horizon (situated beneath the drainage depth) is less permeable with a lower degree of preferential flow occurring through macropores (See Kjær *et al.* 2005c for details).

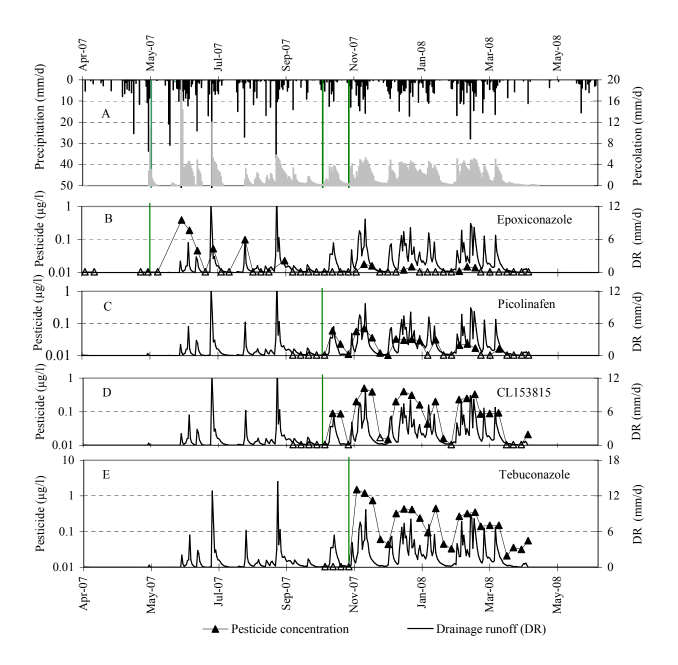


Figure 30. Precipitation and simulated percolation (A) together with concentration of Epoxiconazole (B), Picolinafen (C), CL153815 (D) and Tebuconazole (E) in the drainage runoff (DR on the secondary axis) at Estrup in 2007/2008. The green vertical lines indicate the dates of application. Open triangles indicate values below the detection limit of $0.01 \, \mu g/l$.

6 Pesticide leaching at Faardrup

6.1 Materials and methods

6.1.1 Site description and monitoring design

Faardrup is located in southern Zealand (Figure 1). The test field covers a cultivated area of 2.3 ha (150 x 160 m). The terrain slopes gently to the west by $1-3^{\circ}$ (Figure 31). Based on three profiles in the buffer zone bordering the field, the soil was classified as Haplic Vermudoll, Oxyaquic Hapludoll and Oxyaquic Argiudoll (Soil Survey Staff, 1999). The topsoil is characterized as sandy loam with 14–15% clay and 1.4% organic carbon. Within the upper 1.5 m numerous desiccation cracks coated with clay are present. The test field contains glacial deposits dominated by sandy till to a depth of about 1.5 m overlying a clayey till. The geological description shows that small channels or basins filled with meltwater clay and sand occur both interbedded in the till and as a large structure crossing the test field (Lindhardt et al., 2001). The calcareous matrix and the reduced matrix begin at 1.5 m and 4.2 m b.g.s., respectively. The dominant direction of groundwater flow is towards the west in the upper part of the aquifer (Figure 31). During the monitoring period the groundwater table was located 1– 2 and 2-3 m b.g.s. in the lower and upper parts of the area, respectively. During fieldwork within the 5 m deep test pit it was observed that most of the water entering the pit came from an intensely horizontally fractured zone in the till at a depth of 1.8-2.5 m. The intensely fractured zone could very well be hydraulically connected to the sand fill in the deep channel, which might drain part of the percolation. The bromide tracer study showed, however, that virtually none of the applied bromide reached the vertical monitoring well (M6) located in the sand-filled basin (Figure 32), thus indicating that hydraulic contact with the surface in the "basin" does not differ from that in other parts of the test field, and that the basin is a small pond filled with sediments from local sources. A brief description of the sampling procedure is provided in Appendix 2. The monitoring design and test site are described in detail in Lindhardt et al. (2001) and the analysis methods in Kjær et al. (2002).

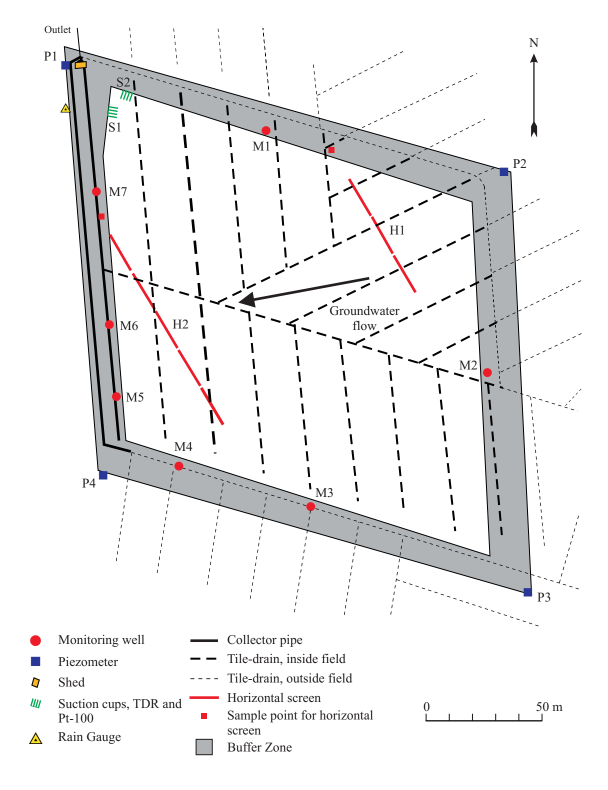


Figure 31. Overview of the Faardrup site. The innermost white area indicates the cultivated land, while the grey area indicates the surrounding buffer zone. The positions of the various installations are indicated, as is the direction of groundwater flow (by an arrow).

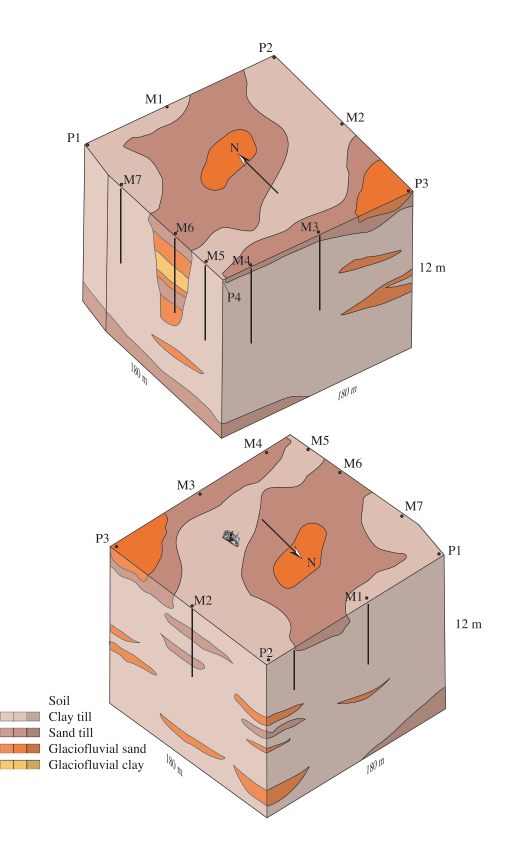


Figure 32. Geological description of Faardrup (Lindhardt et al. 2001).

6.1.2 Agricultural management

Management practice during the two recent growing seasons is briefly summarized below and detailed in Appendix 3 (Table A3.5). For information about management practice during the previous monitoring periods see Kjær *et al.* (2002, 2003, 2004, 2005c, 2007 & 2008).

On 17 August 2006 the field was ploughed and sown with winter rape (cv. Labrador) and the following day sprayed with the herbicide clomazone. This substance was not included in the monitoring, however. The rape was sprayed again on 19 February 2007, when the rape had six unfurled leaves, using the herbicide propyzamide, and again on 30 March when the rape had seven visibly extended internodes, using clopyralid (not included in the monitoring programme). At harvest on 19 July seed yield was 30.0 hkg/ha (91% dry matter). The straw, 68.5 hkg/ha (100% dry matter), was shredded at harvest. Ploughing and sowing of a winter wheat crop (cv. Ambition) was done on 18 September 2007. Ten days later the wheat emerged. Pendimethalin was used against weeds on 9 October when the wheat had two leaves unfurled. The fungicide tebuconazol was applied on 20 November. On 20 August 2008 grain yield amounted to 89.6 hkg/ha (85% dry matter), slightly above the average for the soil type for this year (Plantedirektoratet, 2008). Straw yield was 65.2 hkg/ha (100% dry matter).

6.1.3 Model setup and calibration

The numerical model MACRO (version 5.1) was applied to the Faardrup site covering the soil profile to a depth of 5 m b.g.s., always including the groundwater table. The model was used to simulate the water flow in the unsaturated zone during the full monitoring period September 1999 - June 2008 and to establish an annual water balance.

Compared to the setup in Kjær *et al.* (2008), a year of validation was added to the MACRO setup for the Faardrup site. The setup was accordingly calibrated for the monitoring period May 1999 - June 2004 and validated for the monitoring period July 2004 - June 2008. For this purpose, the following time series were used: observed groundwater table measured in the piezometers located in the buffer zone, water content measured at three depths (25, 60 and 110 cm b.g.s.) from the two profiles S1 and S2 (Figure 35) and measured drainage flow. Data acquisition and model setup are described in Barlebo *et al.* (2007) and Rosenbom *et al.* (in prep.).

Due to electronic problems, precipitation measured at Flakkebjerg located 3 km east of Faardrup was used for the monitoring periods: July 1999 - June 2002, July 2003 - June 2004, January and February of both 2005 and 2006, and July 2006 - June 2007. Precipitation measured locally at Faardrup was used for the rest of the monitoring period.

Table 12. Annual water balance for Faardrup (mm/year). Precipitation is corrected to the soil surface according to the method of Allerup and Madsen (1979).

	Normal precipitation ¹⁾	Precipitation ²⁾	Actual evapotranspiration	Measured drainage	Simulated drainage	Groundwater recharge ³⁾
1.7.99-30.6.00	626	715	572	192	152	-50
1.7.00-30.6.01	626	639	383	50	34	206
1.7.01-30.6.02	626	810	515	197	201	98
1.7.02-30.6.03	626	636	480	49	72	107
1.7.03-30.6.04	626	685	505	36	18	144
1.7.04-30.6.05	626	671	469	131	55	72
1.7.05-30.6.06	626	557	372	28	15	157
1.7.06-30.6.07	626	790	515	202	212	72
1.7.07-30.6.08	626	645	521	111	65	13

¹⁾ Normal values based on time series for 1961–1990.

6.2 Results and discussion

6.2.1 Soil water dynamics and water balances

The level and dynamics of the soil water saturation in all three horizons in the hydraulic year July 2007 - June 2008 were generally well described by the model (Figure 33D, 33E, and 33F). However, for the summer period 2007 the model overestimated the drop in level of the measured groundwater table (Figure 33B). Because of this overestimate, drainage flow was measured but not simulated in December 2007.

The resulting water balance for Faardrup for the nine monitoring periods is shown in Table 12. Compared with the previous eight years, the latest hydraulic year July 2007 - June 2008 was characterised by having the fourth lowest precipitation, the second highest simulated actual evapotranspiration and the fourth highest measured and simulated drainage. Precipitation in this year was characterized by May-June being very dry and July and March being very wet (Appendix 4). Due to this precipitation pattern, the duration of the simulated percolation period of the year July 2007 - June 2008 was represented by continuous percolation throughout the period October-May (Figure 33A). Compared to the other years, the climate this year gave rise to a longer period where the groundwater table was a bit higher than the drainage level, causing a continuous long-term contribution to the drains (Figure 33B and 33C). Apart from the differences in measured and simulated yearly amount of drainage, the overall trend in the drainage pattern of the hydraulic years was captured.

²⁾ For July 1999 - June 2002, July 2003 - June 2004, in January and February of both 2005 and 2006, and July 2006 - June 2007, measured at the DIAS Flakkebjerg meteorological station located 3 km from the test site (see detailed text above).

³⁾ Groundwater recharge is calculated as precipitation minus actual evapotranspiration minus measured drainage.

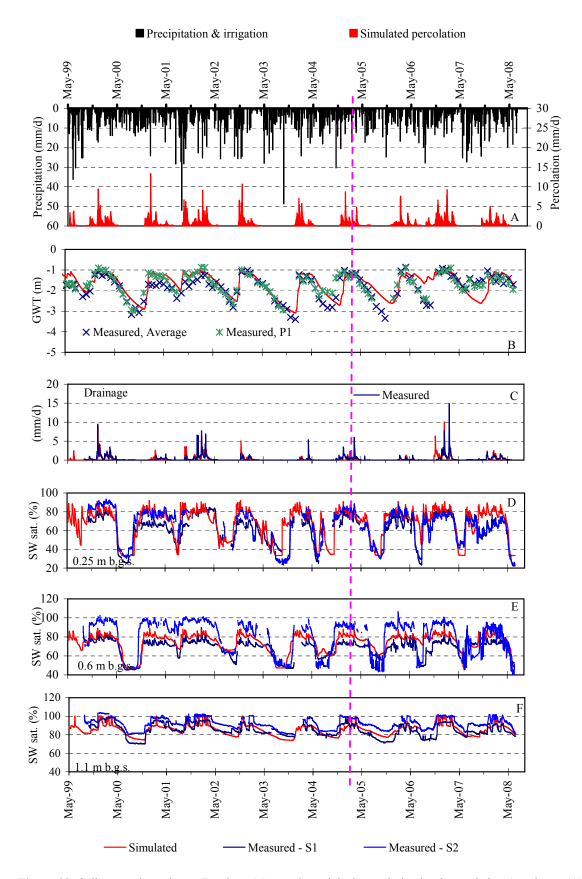


Figure 33. Soil water dynamics at Faardrup: Measured precipitation and simulated percolation 1 m b.g.s. (A), simulated and measured groundwater level GWT (B), simulated and measured drainage flow (C) and simulated and measured soil water saturation (SW sat.) at three different soil depths (D, E, and F). The measured data in B derive from piezometers located in the buffer zone. The measured data in D, E and F derive from TDR probes installed at S1 and S2 (Figure 35). The broken vertical line indicates the beginning of the validation period (July 2004 - June 2008).

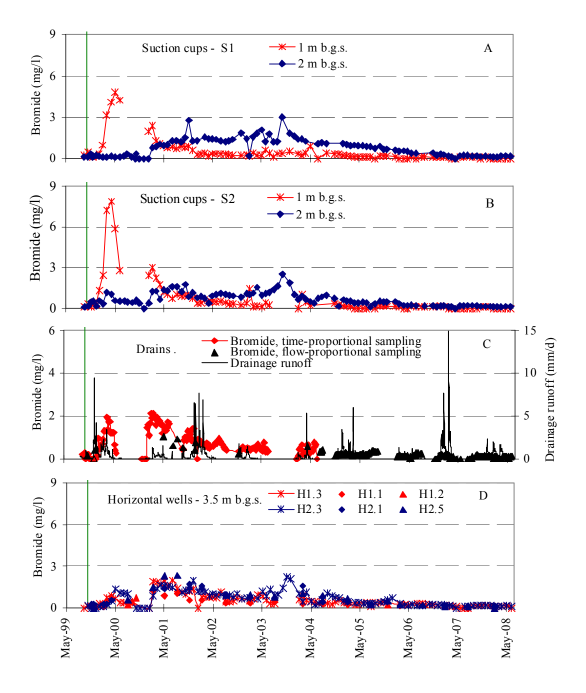


Figure 34. Bromide concentrations at Faardrup. A and B refer to suction cups located at S1 and S2. The bromide concentration is also shown for drainage runoff (C) and the horizontal monitoring wells (D). The green vertical line indicates the date of bromide application.

6.2.2 Bromide leaching

The bromide concentration shown in Figure 34 and Figure 35 relates to the bromide applied in May 2000, as described further in Kjær *et al.* 2003, and further evaluated in Rosenbom et al. (in prep).

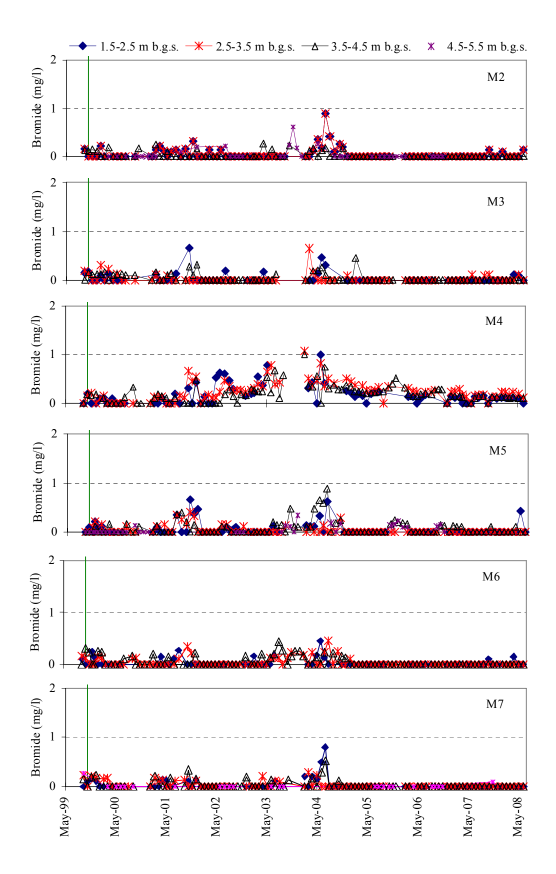


Figure 35. Bromide concentrations at Faardrup. The data derive from the vertical monitoring wells (M2–M7). Screen depth is indicated in m b.g.s. The green vertical line indicates the date of bromide application.

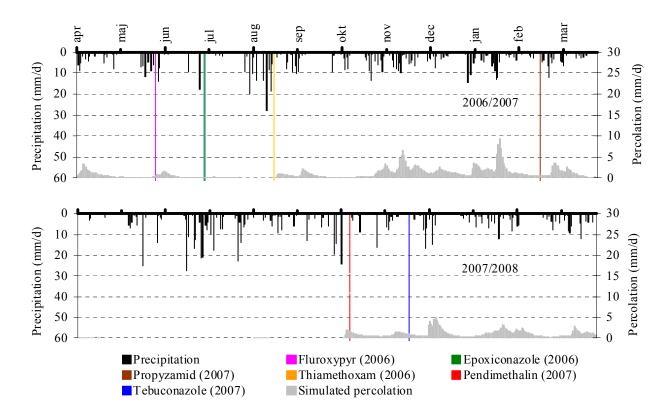


Figure 36. Pesticide application, precipitation (primary axis) together with simulated percolation (secondary axis) at Faardrup in 2006/2007 (upper) 2007/2008 (lower).

6.2.3 Pesticide leaching

Monitoring at Faardrup began in September 1999 and presently encompasses several pesticides and their degradation products, as indicated in Table 13. Pesticide application during the two most recent growing seasons is shown together with precipitation and simulated precipitation in Figure 36. It should be noted that precipitation is corrected to the soil surface according to Allerup and Madsen (1979), whereas percolation (1 m b.g.s.) refers to accumulated values as simulated with the MACRO model. It should also be noted that as tribenuronmethyl (applied here as Express) degrades rapidly, the leaching risk is more associated with its degradation product, triazinamin-methyl. For the same reason it is the degradation product and not the parent compound that is monitored in the PLAP (Table 13).

Table 13. Pesticides analysed at Faardrup with the product used shown in parentheses. Degradation products are in italics. Precipitation (prec.) and percolation (perc.) are accumulated from the date of first application (app. date) until the end of monitoring. 1^{st} month perc. refers to accumulated percolation within the first month after application. C_{mean} refers to average leachate concentration in the drainage water the first drainage season after application (See Appendix 2 for calculation methods). The number of pesticide-positive samples is indicated in parentheses.

Crop and analysed pesticides Application End of Prec. Perc. 1st month C_{mean} date monitoring (mm) (mm) perc. (mm) $(\mu g/l)$ Winter wheat 1999 Glyphosate (Roundup 2000) Aug 99 947 0 $<0.01(8^*)$ Apr 03 2526 $< 0.01(17^*)$ - AMPA Bromoxynil (Briotril) Oct 99 < 0.01(0) Apr 02 1738 751 35 Ioxynil (Briotril) < 0.01(2) Oct 99 Apr 02 1738 751 35 Apr 02 Fluroxypyr (Starane 180) Apr 00 1408 494 < 0.01(2) Propiconazole (Tilt Top) May 00 Jul 03 2151 669 0 $<0.01(1^*)$ Fenpropimorph (Tilt Top) May 00 Jul 02 1518 491 < 0.01(1) 0 - fenpropimorphic acid < 0.01(0) Pirimicarb (Pirimor G) Jun 00 Jul 03 2066 684 0 $< 0.01(9^*)$ < 0.01(9*) - pirimicarb-desmethyl - pirimicarb-desmethyl-formamido $< 0.02(5^*)$ Sugar beet 2001 Glyphosate (Roundup 2000) Oct 00 Jul 03 1747 709 0 < 0.01(8*) - AMPA $0.01(17^*)$ Metamitron (Goltix WG) May 01 Jul 03 1512 507 4 0.01(35) - metamitron-desamino 0.01(63)May 01 507 Jul 03 1512 4 Ethofumesate (Betanal Optima) 0.06(45)< 0.01(0) Desmedipham (Betanal Optima) May 01 Jul 03 1512 507 < 0.02(0) Phenmedipham (Betanal Optima) May 01 Jul 03 1512 507 4 < 0.01(2) - MHPC < 0.02(3) Fluazifop-P-butyl (Fusilade X-tra) Jun 01 Jul 03 0 1460 503 < 0.01(0) - fluazifop (free acid) 0.02(17)<0.01(9*) Pirimicarb (Pirimor G) Jul 01 Jul 03 1460 503 1 - pirimicarb-desmethyl $<0.01(9^*)$ - pirimicarb-desmethyl-formamido $< 0.02(5^*)$ Spring barley 2002 < 0.01(1) Flamprop-M-isopropyl (Barnon Plus 3) May 02 Jul 04 1337 333 0 - Flamprop-M (free acid) < 0.01(1) MCPA (Metaxon) May 02 Jul 04 1358 337 4 $<0.01(2^*)$ - 4-chlor,2-methylphenol $<0.02(1^*)$ - Triazinamin-methyl ¹⁾ (Express) May 02 Jul 04 1358 4 < 0.02(0) 337 Dimethoate (Perfekthion 500 S) Jun 02 Jul 04 1328 333 0 < 0.01(0) Jun 02 Propiconazole (Tilt 250 EC) Jul 04 1328 < 0.01(1*) 333 0

The leaching risk of pesticides applied during 2000, 2001, 2002, 2004 and 2005 has been evaluated in Kjær *et al.* (2002, 2003, 2004, 2005c, 2007 and 2008).

Since the metabolites of terbuthylazine (applied in 2005) and its metabolites were still included in the current monitoring period, the results of these applications are summarised below.

Terbuthylazine (applied in May 2005 on a maize crop), Table 13, Figure 37 and 38 and its metabolite desethyl-terbuthylazine leached both to the drainage system and to groundwater monitoring screens in concentrations exceeding 0.1 μ g/l. Average concentrations in the drainage water the first year after application amounted to 0.67 μ g/l and 0.59 μ g/l, respectively.

Systematic chemical nomenclature for the analysed pesticides is given in Appendix 1.

¹⁾ Degradation product of tribenuron- methyl. The parent compound degrades too rapidly to be detected by monitoring.

^{*} Pesticide has been applied twice, and the findings are not necessarily related to one specific application.

[†] Monitoring will continue during the following year. The values for prec. and perc. are accumulated up to July 2004.

Table 13 continued. Pesticides analysed at Faardrup with the product used shown in parentheses. Degradation products are in italics. Precipitation (prec.) and percolation (perc.) are accumulated from the date of first application (app. date) until the end of monitoring. 1st month perc. refers to accumulated percolation within the first month after application. C_{mean} refers to average leachate concentration in the drainage water the first drainage season after application (See Appendix 2 for calculation methods). The number of pesticide-positive samples is indicated in parentheses.

Crop and analysed pesticides	Application Date	End of monitoring	Prec. (mm)	Perc. (mm)	1 st month perc. (mm)	C _{mean} (µg/l)
Winter rape 2003 Clomazone (Command CS) - propanamide-clomazone (FMC65317)	Aug 02	Apr 05	1761	509	4	<0.02(1) <0.02(1)
Winter wheat 2004						
Prosulfocarb (Boxer EC)	Oct 03	Apr 06	1542	454	0	< 0.01 (0)
MCPA (Metaxon) - 4-chlor,2-methylphenol	Jun 04	Jul 06	1307	331	0	<0.01 (2*) <0.01(1*)
Azoxystrobin (Amistar) - CyPM	Jun 04	Jul 07	2098	636	0	<0.01 (0) <0.01 (4)
Maize 2005						
Terbuthylazine (Inter-Terbutylazin)	May 05	Jul 08	2078	666	4	0.67 (93)
- desethyl-terbuthylazine	May 05	Jul 08	2078	666		0.59 (156)
- 2-hydroxy-terbuthylazine	May 05	Jul 08	2078	666		0.04 (56)
- desisopropyl-atrazine	May 05	Jul 08	2078	666		0.03 (85)
- 2- hydroxy-desethyl-terbuthylazine	May 05	Jul 07	1428	465	4	0.07 (16)
Bentazone (Laddok TE) - AIBA	May 05	Jul 07	1408	464	6	2.82 (28) <0.01 (1)
Spring barley 2006						
Fluroxypyr (Starane 180 S)	May 06	Jul 08	1496	524	17	< 0.02(2)
Epoxiconazole (Opus)	Jun 06	Jul 08	1441	507	3	< 0.01 (0)
Winter Rape 2007						
Thiamethoxam (Cruiser RAPS)	Aug 06	Jul 08	1304	505	27	<0.01 (0)
- CGA 322704	Feb 07	Jul 08 [†]	916	234	46	<0.02 (0)
Propyzamide (Kerb 500 SC) - RH-24644	reb 07	Jul 08	910	234	40	$0.138(5)^{1}$
- КН-24044 - RH-24580						<0.01 (4) ¹⁾ <0.01 (0) ¹⁾
- KH-24560 - RH-24655						<0.01 (0) 1 <0.01 (1) 1)
Winter wheat 2008						
Pendimethalin (Stomp)	Oct 07	Jul 08 [†]	385	201	24	< 0.01 (2)
Tebuconazol (Folicur EC 250)	Nov 07	Jul 08 [†]	329	163	56	<0.01 (5)

Systematic chemical nomenclature for the analysed pesticides is given in Appendix 1.

While leaching of terbuthylazine ceased two years after application (last detected in the drainage water on 28 August 2007), minor root zone leaching of desethyl-terbuthylazine was observed during the entire monitoring period (e.g. four years after application). Terbuthylazine and especially desethyl-terbuthylazine were also found in high concentrations in the groundwater monitoring well M5 (Figure 38B and C). During the first two years after application concentrations of terbuthylazine and especially desethyl-terbuthylazine frequently exceeded 0.1 µg/l in M5. During the last monitoring year, concentrations decreased and all concentrations were below 0.1 µg/l. Minor leaching of the metabolites desisopropyl-atrazine, 2-hydroxy-desethyl-terbuthylazine and 2-hydroxy-terbuthylazine was also observed, but at low frequencies of detection and concentration levels in the drainage water system and groundwater monitoring wells (Figures 37 and 38 and Table A5.5 in Appendix 5).

^{*} Pesticide has been applied twice, and the findings are not necessarily related to one specific application.

[†] Monitoring will continue during the following year. The values for prec. and perc. are accumulated up to July 2004.

1) Drainage runoff commenced prior to the application of propyzamide and the weighted concentrations refer to the period from the date of application (Feb 07) until 1 July 2007.

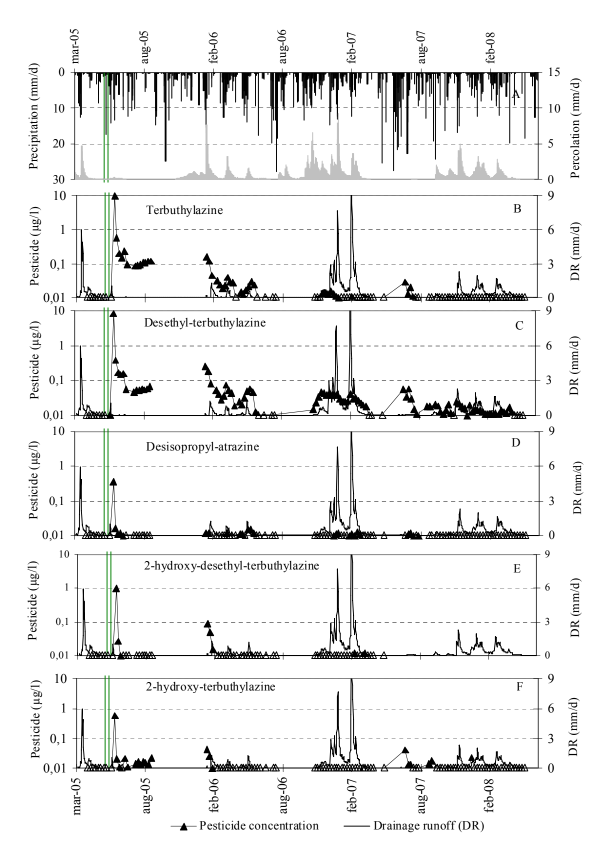


Figure 37. Precipitation and simulated percolation (A) together with concentration of terbuthylazine (B), desethylterbuthylazine (C), 2-hydroxy-terbuthylazine (D), desisopropyl-atrazine (E) and 2-hydroxy-desethyl-terbuthylazine (F) in the drainage runoff (DR on secondary axis) at Faardrup. The green vertical lines indicate the dates of applications. Open triangles indicate values below the detection limit of 0.01 μg/l.

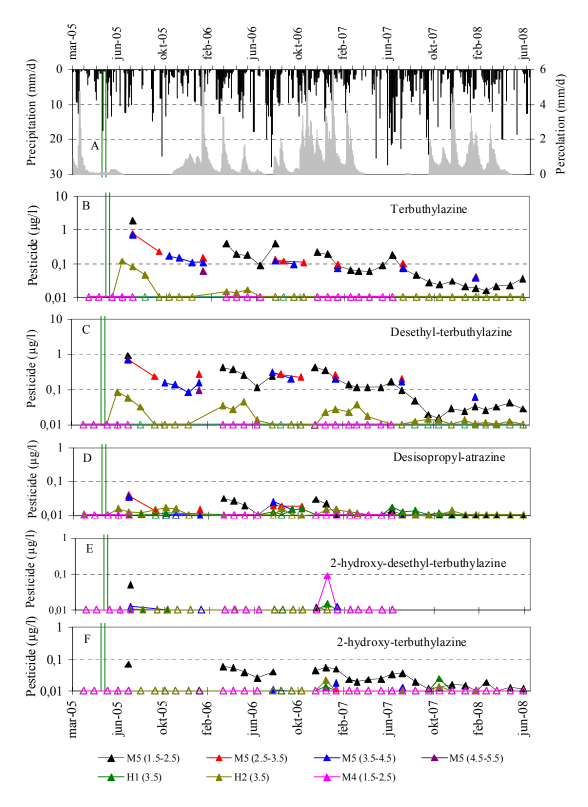


Figure 38. Precipitation and simulated percolation (A) together with concentration of terbuthylazine (B), desethylterbuthylazine (C), desisopropyl-atrazine (D), 2-hydroxy-desethyl-terbuthylazine (E) and 2-hydroxy-terbuthylazine in monitoring well H1, H2, M5 and the uppermost screen of M4 at Faardrup. None of these pesticides were detected in the lower screens of M4 (monitored monthly) nor in M6 (monitored half-yearly). The green vertical lines indicate the dates of applications, numbers in parentheses screen depth and open triangles values below the detection limit of $0.01 \, \mu g/l$.

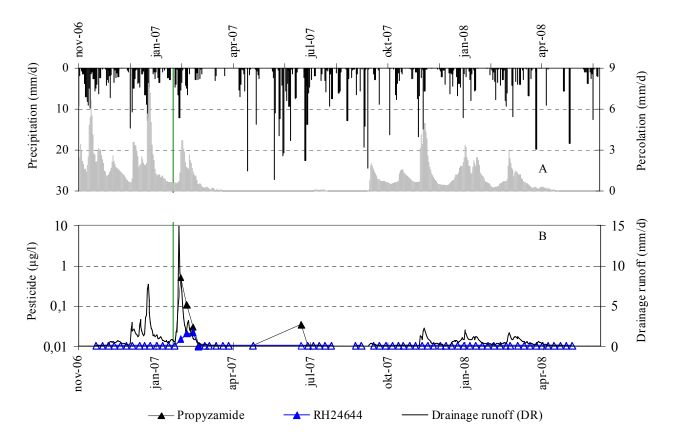


Figure 39. Precipitation and simulated percolation (A) together with concentration of propyzamide and RH24644 (B) in the drainage runoff (DR on secondary axis) at Faardrup in 2006/2007. The green vertical lines indicate the date of applications. Open triangles indicate values below the detection limit of 0.01 µg/l.

Fluroxypyr and epoxiconazole were applied in May and June 2006. Both pesticides have been monitored until July 2008. Only fluroxypyr has been found, but in low concentrations and only in two samples (Table 13).

Propyzamide and one of its metabolites leached to the drainage system (Figure 39, Table 13). Due to four very high concentrations of propyzamide, the calculated average concentration amounted to 0.138 μ g/l. It should be noted that drainage runoff commenced more than two and a half months prior to the application of propyzamide and the weighted concentrations refer to a period of 4.5 months (17 February - 1 July 2007). Figure 38 shows that propyzamide and the metabolite RH24644 were only found in high concentrations in the first flow event following the application, and that none of them have been detected since July 2007. Propyzamide was detected once in one of the horizontal screens 3.5 metre b.g.s. at a concentration of 0.033 μ g/l (data not shown).

The leaching risk of pesticides applied in 2007 will not be evaluated until the 2008 monitoring results become available, i.e. when two years of monitoring data have been collected. Tebuconazole and pendimethalin have until now been detected in five and two samples, respectively. However, measured concentrations never exceeded 0.1 µg/l.

7 Pesticide analysis quality assurance

Reliable results and scientifically valid methods of analysis are essential for the integrity of the present monitoring programme. Consequently, the field monitoring work has been supported by intensive quality assurance entailing continuous evaluation of the analyses employed. Two types of sample are used in the quality control – samples with known pesticide composition and concentration are used for *internal monitoring* of the laboratory method, while *external spiked samples* are used to incorporate additional procedures such as sample handling, transport and storage. Pesticide analysis quality assurance (QA) data for the period July 2007 - June 2008 are presented below, while those for the preceding monitoring periods are given in Kjær *et al.* (2002, 2003, 2004, 2005c, 2007 and 2008).

7.1 Materials and methods

Apart from chlormequat, which was analysed at GEUS, all pesticide analyses were carried out at commercial laboratories selected on the basis of a competitive tender. In order to assure the quality of the analyses, the call for tenders included requirements as to the laboratory's quality assurance (QA) system comprising both an internal and an external control procedure. In addition to specific quality control under the PLAP, the laboratory takes part in the proficiency test scheme employed by the Danish Environmental Protection Agency when approving laboratories for the Nationwide Monitoring and Assessment Programme for the Aquatic and Terrestrial Environments (NOVANA).

7.1.1 Internal QA

With each batch of samples the laboratory analysed one or two control samples prepared at each laboratory as part of their standard method of analysis. The pesticide concentration in the internal QA samples ranged between 0.05–0.06 μ g/l. When using these data it was possible to calculate and separate the analytical standard deviation into within-day (S_w), between-day (S_b) and total standard deviation (S_t). Total standard deviation was calculated using the following formula (Wilson 1970, Danish EPA 1997):

$$S_t = \sqrt{S_w^2 + S_b^2}$$

7.1.2 External QA

Every four months, two external control samples were analysed at the laboratories along with the various water samples from the five test sites. Two stock solutions of different concentrations were prepared from two standard mixtures in ampoules prepared by Dr. Ehrenstorfer, Germany (Table 14). Fresh ampoules were used for each set of standard solutions. For preparation of the controls the pesticide mixture was pipetted into a preparation glass containing 10 ml of ultrapure water. The glass was closed and shaken thoroughly and shipped to the staff collecting the samples. The staff finished the preparation of control samples in the field by quantitatively transferring the standard

solution to a 3 l measuring flask. The standard solution was diluted and adjusted to the mark with groundwater from an upstream well. After thorough mixing, the control sample was transferred to a sample bottle and transported to the laboratories together with the regular samples. The standard solutions were prepared two days before a sampling day. Chlormequat was not included in the ampoules, and the standard solution was therefore made directly from the analytic standard and subsequently transferred into a 1 l measuring flask in the field following the same procedure as for the other pesticides.

The pesticide concentration in the solution is indicated in Table 14. Blank samples consisting of HPLC water were also included in the external QA procedure every month. All samples included in the control were labelled with coded reference numbers so that the laboratories were unaware of which samples were controls and which were blanks.

Table 14. Pesticide concentrations in both the original ampoules and in the resulting high-level and low-level

external control samples

Compound	Original ampoules	High-level control	Low-level control
	(mg/l)	$(\mu g/l)$	$(\mu g/l)$
AMPA	1.0	0.117	0.050
Bentazone	1.0	0.117	0.050
CL153815	1.0	0.117	0.050
Chlormequat*		No control	0.039
CyPM	1.2	0.140	0.078
Epoxiconazole	1.0	0.117	0.050
Florasulam	1.0	0.117	0.050
Fluroxypyr	1.0	0.117	0.050
Glyphosate	1.0	0.117	0.050
Metsulfuron-methyl	1.0	0.117	0.050
Pendimethalin	1.2	0.140	0.078
PPU (IN70941)	1.0	0.117	0.050
Propyzamide	1.0	0.117	0.050
Tebuconazole	1.0	0.117	0.050
Terbuthylazine-desethyl	1.0	0.117	0.050
Thiamethoxam	1.0	0.117	0.050

^{*} Chlormequat was not included in the ampoules and the standard solution was therefore made directly from the analytic standard

7.2 Results and discussion

7.2.1 Internal OA

Ideally, the analytical procedure should provide precise and accurate results. However, the pesticide analyses are subject to a certain standard deviation. Such standard deviation may be the combined result of several contributing factors. Overall, the accuracy of an analytical result reflects two types of error: Random errors related to precision and systematic errors relating to bias. In a programme like PLAP it is relevant to consider possible changes in analytical "reliability over time". As these errors may change over time it is relevant to distinguish between standard deviations resulting from within-day variation as opposed to those associated with between-day variation in the analytical result. To this end, control samples are included in the analytical process as described above. Thus, by means of statistical analysis of the internal QA data it is possible to separate and estimate the different causes of the analytical variation in two categories: day-to-day variation and within-day variation (Miller et al., 2000; Funk et al., 1995). This kind of analysis can provide an indication of the reliability of the

analytical results used in the PLAP. The statistical tool used is an analysis of variance (ANOVA) and encompasses all duplicate pesticide analyses, single analyses being excluded. The analysis can be divided into three stages:

- 1. **Normality**: An initial test for normality is made as this is an underlying assumption for the one-way ANOVA.
- 2. **Between-day contribution:** Explained simply, this test will reveal any day-to-day contribution to the variance in the measurements. If there is none, the total standard deviation can be considered to be attributable to the within-day error of the analysis. For this purpose an ANOVA-based test is used to determine if the between-day standard deviation (S_b) differs significantly from 0 (this test is made as an F-test with the H₀: between-day mean square = within-day mean square).
- 3. Calculating standard deviations: If the F-test described above reveals a contribution from the between-day standard deviation (S_b) , it is relevant to calculate three values: The within-day standard deviation S_w , the between-day standard deviation S_b , and the total standard deviation S_t .

As the error associated with the analytical result is likely to be highly dependent on the compound analysed, the QA applied is pesticide-specific. The results of the internal QA statistical analysis for each pesticide are presented in Table 15. For reference, estimated S_b values are listed for all pesticides, including those for which the between-day variance is not significantly greater than the within-day variance. ANOVA details and variance estimates are also included, even for pesticides where the requirement for normality is not fulfilled. Such data should obviously be interpreted with caution.

As a "rule of thumb", the between-day standard deviation should be no more than double the within-day standard deviation. From Table 15 it can be seen that S_b/S_w ratios greater than two were observed for several compounds. For three compounds, the results indicate that day-to-day variation makes a significant contribution. Among the compounds meeting the normality requirement, the S_b/S_w ratio is highest for CL153815 mesosulfuron-methyl and tebuconazole. When all compounds are considered, a relatively high S_b/S_w ratio is apparent for propyzamide and tebuconazole. This relatively high values are caused by very low within-day standard deviation, i.e. within each laboratory day, the variation on the propyzamide and tebuconazole analysis is small compared to the other compounds, whereas the variation between days is comparable to the other compounds analysed. Thus, low values of S_w rather than critical values of S_b caused the high ratios, as reflected by the low S_t . As reflected by the data in Table 15, the four compounds with significant between-day contribution were AMPA, florasulam, glyphosate, and metribuzin-diketo.

The total standard deviations (S_t) of the various analyses of pesticides and degradation products lie within the range 0.001-0.016 μ g/l (only data with n≥3 are included in the analysis). The overall mean S_t was 0.007 μ g/l.

Table 15. Internal QA of pesticide analyses carried out in the periode 1.7.2007-30.6.08. Results of the test for normality, one-way analysis of variance (ANOVA), the estimated values of standard deviations (w: within-day, b: between-day, t: total – see text for details), pesticide concentration in internal QA sample (Conc.) and number of duplicate samples (n) are given for each pesticide. Degradation products are indicated with 'D' following the compound name. For tests the P value α =0.05 was used. Only data for n≥3 are included.

Pesticide/Degradation product	Normal	Significant Sb	Sw	Sb	St	Ratio	N	Conc
	distribution α =0.05	Between day contribution	$(\mu g/l)$	$(\mu g/l)$	$(\mu g/l)$	Sb/Sw		$(\mu g/l)$
	α=0.05	ANOVA						
		α =0.05						
2-hydroxy-desethyl-terbuthylazine (D)			0.008	0.006	0.010	0.77	42	0.050
2-hydroxy-terbuthylazin (D)			0.003	0.004	0.005	1.52	53	0.052
AMPA (D)			0.001	0.000	0.001	0.00	12	0.030
PPU (IN70941) (D)			0.006	0.011	0.013	1.89	23	0.053
PPU-desamino (IN70942) (D)	Yes		0.002	0.006	0.006	2.48	24	0.053
Azoxystrobin			0.003	0.003	0.004	1.18	49	0.053
Bentazone			0.002	0.007	0.007	3.31	43	0.053
CL153815 (D)	Yes		0.003	0.009	0.010	3.13	28	0.129
Clopyralid			0.009	0.013	0.016	1.50	15	0.050
Desethylterbuthylazine (D)			0.002	0.003	0.004	1.62	55	0.053
Desisopropylatrazin (D)			0.003	0.007	0.007	2.60	55	0.053
Epoxiconazole	Yes	Yes	0.003	0.003	0.004	0.88	6	0.050
Florasulam			0.002	0.002	0.003	0.84	45	0.050
Florasulam-desmethyl (D)			0.004	0.005	0.007	1.25	8	0.098
Florasulam-desmethyl (D)			0.004	0.005	0.007	1.25	24	0.117
Glyphosate	Yes		0.001	0.002	0.002	1.41	12	0.030
Mesosulfuron (D)	Yes		0.004	0.010	0.011	2.42	4	0.112
Mesosulfuron (D)	Yes		0.004	0.010	0.011	2.42	4	0.110
Mesosulfuron-methyl	Yes		0.003	0.008	0.009	2.94	4	0.050
Metribuzine-desamino-diketo (D)		Yes	0.008	0.004	0.009	0.51	13	0.050
Metribuzine-diketo (D)		Yes	0.003	0.002	0.004	0.83	13	0.050
Pendimethalin			0.006	0.005	0.008	0.79	50	0.050
Picolinafen	Yes		0.008	0.014	0.016	1.67	28	0.114
Propyzamide			0.001	0.004	0.005	3.48	52	0.053
Tebuconazole	Yes		0.002	0.006	0.006	3.81	6	0.050
Terbuthylazine			0.002	0.003	0.004	1.93	55	0.050
Triazinamin			0.004	0.010	0.011	2.31	19	0.110
Triazinamin-methyl (D)			0.005	0.009	0.010	1.90	13	0.054

7.2.2 External QA

Table 16 provides an overview of the recovery of all externally spiked samples. As the results for each field site in Table 16 are based on only one to three observations for each concentration level (high/low), this should not be interpreted too rigorously. Recovery of the most spiked samples is generally good, averages ranging from 63-111%. The only exception is the very low recovery of glyphosate (average recovery only 29%) and the high recovery of PPU found at Jyndevd (177-247%).

Table 16. External spiked samples. Average recovery (%) of the nominal concentration at low/high concentration level indicated for each site. n_{low} and n_{high} refer to the total number of samples being spiked at low and high concentrations, respectively

	Tyl	strup	Jynd	evad	Sils	trup	Est	rup	Faar	drup	Average	n_{low}/n_{high}
	Low	High	Low	High	Low	High	Low	High	Low	High		
AMPA							63	71			67	3/3
Bentazone							107	105			106	3/3
CL153815							91	83			87	3/3
Cq			73		66		54				64	7/0
CyPM	90	107					71	83			88	4/4
Epoxiconazole	103	111	109	114	94	103	100	105	105	108	105	15/14
Florasulam			100	97			90	91			94	6/6
Fluroxypyr									98	94	96	3/3
Glyphosate							31	27			29	3/3
Metsulfuron												
methyl			99	105							102	3/3
Pendimethalin	68	87			63	86	62	76	65	74	73	10/9
PPU (In70941)	100*	93*	247*	177*							154*	6/5
Propyzamide	95	90			94	94			171	125	112	7/6
Tebuconazole	104	111	112	114			88	96	101	100	103	12/11
Terbuthylazine-												
desethyl							109	114	115	123	115	6/6
Thiamethoxam	107	115							109	114	111	6/5

^{*}Indicates that a degradation product PPU-desamino of the compound was detected even if it was not included in the spiking solution.

As the recovery of glyphosate is markedly lower than the previous year, where annual average recoveries ranged from 71-93% during 2001-2006 and 69% in 2007, it may indicate a possible recovery problem occurring during 2008. Since August 2005 PPU was frequently detected in the upstream well at Jyndevad from which water was collected for the spiked samples (Figure 14). This background level may explain the high recoveries of PPU at Jyndevad (177-247%) (see Table 16). However, except from PPU at Jyndevad no other contamination is observed in the upstream samples used for spiking.

The degradation product PPU-desamido was observed in several samples, even though this substance is not included in the spiked solution (indicated by asterisks in Table 16). As PPU-desamido was not detected in the blank sample matrix used for spiking, its presence in the spiked samples is likely to derive from the added PPU, the primary degradation product of rimsulfuron degradation. The secondary degradation product PPU-desamido was observed in all the samples spiked with PPU at high-level, and also in 33% (Tylstrup) and 43% (Jyndevad) of the low-level spiked samples. The measured concentration of PPU-desamido at Tylstrup was 28-47% (on average 37%) of the added amount of PPU for the period 2004-2008. The measured concentration of PPUdesamido at Jyndevad was 22-44% (on average 31%) of the added amount of PPU for the period 2003-2005. Due to the presence of a background level of PPU in the groundwater at Jyndevad since August 2005, later data were not included. The concentration of PPU detected at Tylstrup and Jyndevad (Figure 7 and Figure 13) may, therefore, to some extent be underestimated. Finally, the estimated amounts of PPU degraded during transport and storage are similar to those reported earlier (Kjær et al. 2007).

Contamination of samples during collection, storage and analysis is not likely to occur. In a total of 99 blank samples no pesticides or degradation product were detected.

During the 2007/2008 monitoring period a total of 11 pesticides and nine degradation products were detected in samples from the experimental fields, and the external and internal QA data relating to these particular pesticides/degradation products are of special interest. These data (when available) are therefore illustrated in Appendix 6.

7.3 Summary and concluding remarks

The overall quality of the pesticide analysis was considered satisfactory. The QA system showed that:

- Reproducibility of the pesticide analyses was good, total standard deviation being in the range 0.001- $0.016 \mu g/l$.
- Recovery was generally good (average recovery ranging between 67–112%) in external spiked samples. Low recovery of glyphosate was, however, observed in all six samples.
- Contamination of samples during collection, storage and analysis is not likely to occur. In a total of 99 blank samples no pesticides or pesticide degradation products were detected.

8 Summary of monitoring results

This section summarizes the monitoring data from the entire monitoring period, i.e. both data from the two most recent monitoring years (detailed in this report) and data from the previous monitoring years (detailed in previous reports Kjær *et al.*, 2002, 2003, 2004, 2005c, 2007 and 2008). Pesticide detections in samples from the drainage systems, suction cups and monitoring wells are detailed in Appendix 5. The monitoring data reveal that the applied pesticides exhibit three different leaching patterns – no leaching, slight leaching and pronounced leaching (see Table 17). Pronounced leaching is here defined as root zone leaching (1 m b.g.s.) exceeding an average concentration of 0.1 μ g/l. On sandy and loamy soils, leaching is determined as the weighted average concentration in soil water and drainage water, respectively (See Appendix 2). It should be noted, though, that the present evaluation of the leaching risk of some of these pesticides is still preliminary as their potential leaching period extends beyond the current monitoring period. This applies to those pesticides marked with a single asterisk in Table 17. Twelve of the applied pesticides or their degradation products exhibited pronounced leaching.

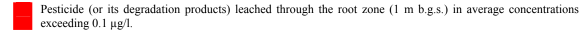
- Two degradation products of metribuzin metribuzin-diketo and metribuzin-desamino-diketo leached from the root zone (1 m b.g.s.) at average concentrations exceeding 0.1 μg/l in the sandy soil at Tylstrup. Both degradation products appear to be relatively stable and leached for a long period of time. Average concentrations reaching 0.1 μg/l were thus seen as long as three years after application. Evidence was also found that their degradation products might be present in the groundwater several years after application. At both sandy sites (Tylstrup and Jyndevad), previous application of metribuzin has caused marked groundwater contamination with its degradation products (see Kjær *et al.*, 2005b for details).
- Glyphosate was found to leach through the root zone at high average concentrations on loamy soils. At the loamy sites Estrup and Silstrup, glyphosate leached from the root zone into the drainage water at average concentrations exceeding 0.1 µg/l. At the Estrup site its degradation product AMPA leached at an average concentration exceeding 0.1 µg/l. The leaching of glyphosate was mainly governed by pronounced macropore flow occurring within the first months after application. AMPA was frequently detected as long as two years after application. That leaching of AMPA occurs for such a relatively long period after application may indicate that AMPA is retained within the soil and gradually released over a very long time, or that glyphosate is retained within the soil and then gradually degraded. So far, marked leaching of AMPA and glyphosate has mainly been confined to the depth of the drainage system and neither has rarely been detected in monitoring screens located below the depth of the drainage system. Glyphosate and AMPA were also detected in drainage water at the loamy site of Faardrup (as well as at the now discontinued Slaeggerup site), but in low concentrations (Kjær et al., 2004). Evidence of glyphosate leaching was only seen on loamy soils, whereas the leaching risk was negligible on the coarse, sandy soil of Jyndevad. Here, infiltrating water passed through a matrix rich in aluminium and iron, thereby providing good conditions for sorption and degradation (see Kjær et al., 2005a for details).

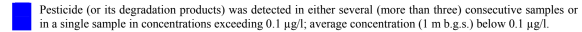
Table 17. Leaching of pesticides or their degradation products at the five PLAP sites. An asterisk indicates pesticides that have been included in the monitoring programme for less than two years. The colours indicate the degree of leaching and the letters H, F, I and GR indicate the type of pesticide: herbicide, fungicide, insecticide and growth retardant, respectively. Pesticides applied in spring 2008 are not included in the table.

	Tylstrup	Jyndevad	Silstrup	Estrup	Faardrup
1: (7)	(Sandy soil)	(Sandy soil)	(Loamy soil)	(Loamy soil)	(Loamy soil)
Azoxystrobin (F)					
Bentazone (H)					
Ethofumesate (H)					
Glyphosate (H)					
Metamitron (H)					
Metribuzine (H)		1)			
Picolinafen (H)		*		*	
Pirimicarb (I)					
Propyzamide (H)					
Rimsulfuron (H)					
Terbuthylazine (H)					
Tebuconazole (F)	*	*		*	*
Amidosulfuron (H)		2)		2)	
Bromoxynil (H)					
Clomazone (H)					
Dimethoate (I)					
Mancozeb (F)					-
Epoxiconazole (F)			*	*	
Fenpropimorph (F)	-		-		
Flamprop-M-isopropyl (H)					
Fluazifop-P-butyl (H)					
Fluroxypyr (H)	-				-
Ioxynil (H)					
MCPA (H)					
Mesosulfuron-methyl (H)					
Pendimethalin (H)					*
Phenmedipham (H)					
Propiconazole (F)					
Prosulfocarb (H)					
Pyridate (H)			-		
Clopyralid (H)	*				
Chlormequat (GR)		*	*	*	
Desmedipham (H)					
Florasulam (H)					
Iodosulfuron-methyl-sodium (H)			*		
			· ·		
Linuron (H) Matsulfuran mathyl (H)					ı
Metsulfuron-methyl (H)					
Thiamethoxam (I)					
Tribenuron-methyl (H)					
Triasulfuron (H) Derived from previous app	1	imr at al. 2002)			

Derived from previous application (see Kjær *et al.*, 2002).

²⁾ Degradation products are not monitored (see text).





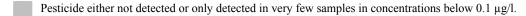


Table 18. Number of samples in which the various pesticides were detected at each site with the maximum concentration $(\mu g/l)$ in parentheses. Degradation products are indicated in italics. The table only encompasses those pesticides/degradation products detected in either several (more than three) consecutive samples or in a single sample

in concentrations exceeding 0.1 µg/l. Pesticides applied in spring 2008 are not included.

in concentrations exceeding 0.1 µg/1.1					P 1
	Tylstrup	Jyndevad	Silstrup	Estrup	Faardrup
Azoxystrobin		0	6(0.034)	65(1.4)	0
-CyPM		0	62(0.34)	132(0.77)	4(0.059)
Bentazone	1(0.012)	30(1.6)	52(6.4)	139(20)	28(43)
- AIBA	0	2(0.034)	0	2(0.06)	1(0.057)
Ethofumesate			24(0.227)	35(3.362)	45(12)
Glyphosate		0	71(4.7)	260(31)	8(0.093)
- AMPA		3(0.022)	137(0.35)	316(1.6)	17(0.11)
Metamitron			69(0.551)	42(26.369)	35(1.7)
- metamitron-desamino			61(0.67)	49(5.549)	63(2.5)
Metribuzine	3(0.024)	0			
- metribuzin-desamino-diketo	317(2.1)	20(1.831)			
- metribuzin-diketo	663(0.69)	29(1.372)			
Picolinafen		0		0	
- CL153815		0		21(0.5)	
Pirimicarb	0	0	17(0.054)	40(0.077)	9(0.056)
- pirimicarb-desmethyl	0	1(0.011)	1(0.052)	0	9(0.053)
- pirimicarb-desmethyl-formamido	0	0	0	26(0.379)	5(0.076)
Propyzamide	0		32(1.6)	, ,	5(0.51)
- RH-24644	0		17(0.051)		4(0.022)
- RH-24580	0		2(0.016)		0
- RH-24655	0		0		1(0.017)
- PPU ⁴	69(0.15)	294(0.29)			-(***-*)
- PPU-desamido ⁴	29(0.042)	137(0.13)			
Terbuthylazine	0	0	96(1.55)	77(11)	93(10)
- desethyl-terbuthylazine	2(0.012)	42(0.056)	269(1.08)	127(8.2)	156(8.3)
- desisopropyl-atrazine	18(0.042)	12(0.050)	47(0.047)*	97(0.44)	85(0.36)
- 2hydroxy- desethyl -terbuthylazine	6(0.026)		29(0.11)*	86(6.3)	16(1)
- 2-hydroxy-terbuthylazine	1(0.04)		26(0.039)*	87(0.99)	56(0.58)
Tebuconazole	1(0.01)	0	20(0.037)	23(1.5)	5(0.045)
Amidosulfuron	1(0.011)	3(0.11)		0	3(0.013)
Bromoxynil	0	0		3(0.6)	0
Clomazone	0	V		3(0.0)	1(0.28)
-propanamide-clomazone	0				1(0.28)
Dimethoate	0	0	2(1.417)	0	0
Epoxiconazole	0	0	0		0
- ETU ^l	9(0.038)	U	U	13(0.39)	U
		2(0,020)	0	1(0.01)	1(0.015)
Fenpropimorph	0	2(0.038)	0	1(0.01)	1(0.015)
- fenpropimorph-acid	0	0	1(0.019)	0	0
Flamprop-M-isopropyl	0		13(0.109)	20(0.069)	1(0.037)
- flamprop (free acid)	0		7(0.096)	13(0.031)	1(0.089)
Fluazifop-P (free acid) ²	0	0	1(0.072)	4(0.050)	17(3.8)
Fluroxypyr	0	0	0	4(0.058)	2(0.19)
Ioxynil	0	0		20(0.25)	2(0.011)
Mesosulfuron-methyl		0		13(0.059)	
MCPA		0	0	12(3.894)	2(0.28)
- 4-chlor-2-methylphenol		0	0	1(0.046)	1(0.24)
Pendimethalin	0	0	14(0.064)		2(0.041)
- MHPC			0		3(0.19)
Phenmedipham			0		2(0.025)
PHCP ³		0	18(2.69)		
Propiconazole	0	0	6(0.033)	27(0.862)	1(0.035)
Prosulfocarb			6(0.18)		0

¹⁾Degradation product of mancozeb. ²⁾Degradation product of fluazifop-P-butyl ³⁾Degradation product of pyridate. ⁴⁾Degradation product of rimsulfuron.

^{*)}Included in the monitoring at Silstrup from February 2003, eight months after application of terbuthylazine.

- Terbuthylazine as well as its degradation products leached through the root zone at high average concentrations on loamy soils. At the three loamy soil sites Silstrup, Estrup and Faardrup, desethyl-terbuthylazine leached from the root zone entering the drainage water in average concentrations exceeding 0.1 µg/l. At the Silstrup (Kjær et al., 2007) and Faardrup sites desethyl-terbuthylazine was frequently detected in the monitoring screens situated beneath the drainage system, concentrations exceeding 0.1 µg/l being detected during a 2- and 24-month period, respectively. Leaching at Estrup (Kjær et al., 2007) was confined to the drainage depth, however. Minor leaching of desethyl-terbuthylazine was also seen at the two sandy sites Jyndevad and Tylstrup, where desethyl-terbuthylazine was detected in low concentrations (<0.1 μg/l) in the soil water sampled 1 m b.g.s. While desethyl-terbuthylazine was not detected in the groundwater monitoring screens at Tylstrup, it was frequently detected in low concentration (< 0.1 µg/l) at Jyndevad (see Kjær et al., 2004). Marked leaching of terbuthylazine was also seen at two of the three loamy sites (Estrup and Faardrup), the leaching pattern being similar to that of desethylterbuthylazine. 2-hydroxy-desethyl-terbuthylazine and 2-hydroxy-terbuthylazine leached at both Faardrup and Estrup and at the latter site the average drainage concentration exceeded 0.1 µg/L. Leaching of these two metabolites was at both sites confined to the drainage system. These two metabolites were not detected in groundwater monitoring screen at Estrup, whereas at Faardrup they were found, however, frequency of detection as well as concentration level being very low.
- One degradation product of rimsulfuron PPU leached from the root zone (1 m b.g.s.) in average concentrations reaching 0.10–0.13 μg/l at the sandy soil site at Jyndevad. PPU appeared to be relatively stable and leached for a long period of time. Average concentrations reaching 0.1 μg/l were thus seen as much as three years after application. With an overall transport time of about four years PPU also reached the downstream monitoring screens. Although the concentration decreased during the last year of monitoring, elevated concentrations in both suction cups and monitoring wells could be found towards the end of the current monitoring period (i.e. six years after application). Minor leaching of PPU was also seen at the sandy site Tylstrup, where low concentrations (0.021 0.11 μg/l) were detected in the soil water sampled 1 and 2 m b.g.s., but only in one groundwater sample from a monitoring well (Appendix 5). It should be noted that the concentration of PPU is likely to be underestimated by up to 22-47%. Results from the field-spiked samples thus indicate that PPU is unstable and may have further degraded to PPU-desamido during subsequent storage and transport.
- In the loamy soil of Estrup ethofumesate, metamitron and its degradation product metamitron-desamino leached through the root zone into the drainage water, average concentrations exceeding 0.1 μg/l. The compounds have not been detected in deeper monitoring screens. These compounds also leached from the root zone at the Silstrup and Faardrup sites, reaching both the drainage system and groundwater monitoring screens. Average concentrations in drainage water were not as high as at Estrup, although concentrations exceeding 0.1 μg/l were observed in both drainage water and groundwater monitoring screens during a 1–6 month period (see Kjær *et al.*, 2002 and Kjær *et al.*, 2004 for details).
- Pirimicarb-desmethyl-formamido (degradation product of pirimicarb) leached through the root zone in high average concentrations from the loamy soil of Estrup

into the drainage water, average concentrations exceeding $0.1~\mu g/l$. Pirimicarb-desmethyl-formamido has not been detected in deeper monitoring screens. Comparable high levels of leaching of pirimicarb-desmethyl-formamido have not been observed with any of the previous applications of pirimicarb at the other PLAP sites (Table 17 and Kjær *et al.*, 2004).

- Bentazone leached through the root zone in average concentrations exceeding 0.1 µg/l in the drainage system at the loamy sites of Silstrup, Estrup and Faardrup. Moreover, bentazone was frequently detected in the monitoring screens situated beneath the drainage system of the Silstrup and Faardrup sites. Apart from a few samples, however, concentrations detected were all below 0.1 µg/l. At Estrup leaching was mostly confined to the depth of the drainage system and rarely detected in deeper monitoring screens (Appendix 5). On the sandy soils bentazone leached at Jyndevad but not at Tylstrup. At Jyndevad, moreover, high concentrations (exceeding 0.1 µg/l) were detected in the soil water samples from suction cups 1 m b.g.s. four months after application. Thereafter, leaching diminished and bentazone was not subsequently detected in the monitoring wells. Although leached in high average concentrations (>0.1 µg/l) at four sites, the leaching of bentazone was generally observed within a short period of time. Initial concentrations of bentazone were usually very high, but then decreased rapidly. In general, concentrations exceeding 0.1 were only found within a period of one to four months following the application.
- Propyzamide leached from the root zone at the loamy Silstrup and Faardrup site, entering the drainage water at average concentrations exceeding 0.1 μg/l Propyzamide was also detected in the monitoring screen situated beneath the drainage system. Apart from a few samples, the concentrations in the screens were always less than 0.1 μg/l, however (Appendix 5).
- Azoxystrobin, and in particular its degradation product CyPM, leached through the root zone in high average concentrations at the loamy sites Silstrup and Estrup. CyPM leached into the drainage water in average concentrations exceeding 0.1 μg/l at both the Silstrup and Estrup sites, while azoxystrobin only leached in concentrations exceeding 0.1 μg/l at the Estrup site. Leaching of azoxystrobin and CyPM has hitherto been confined to the depth of the drainage system, and they have rarely been detected in monitoring screens situated below drainage depth. Neither the loamy Faardrup site nor the sandy Jyndevad site had similar high levels of leaching (Appendix 5).
- CL153815 (degradation product of picolinafen) leached through the root zone in high average concentrations from the loamy soil of Estrup into the drainage water, average concentrations exceeding 0.1 μg/l. CL153815 has not been detected in deeper monitoring screens. Comparable high levels of leaching of CL153815 have not been observed at the sandy soil at Jyndevad, where CL153815 was not detected in any of the collected water samples (Table 18 and Appendix 5).
- Tebuconazole leached through the root zone in high average concentrations from the loamy soil of Estrup into the drainage water, average concentrations exceeding 0.1 µg/l. Leaching of tebuconazole has hitherto been confined to the depth of the drainage system, and it has not been detected in monitoring screens situated

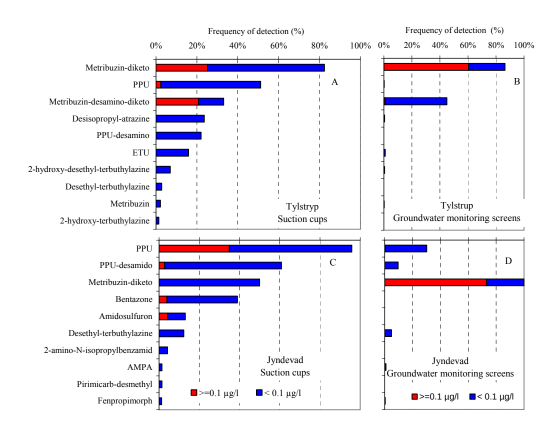


Figure 40. Frequency of detection in samples from the suction cups (left) and groundwater monitoring screens located deeper than the suction cups (right) at the sandy soil sites at Jyndevad (C, D). Frequency is estimated for the entire monitoring period and the time that the different pesticides have been included in the programme and the number of analysed samples thus varies considerably among the different pesticides. The figure only includes the ten most frequently detected pesticides. Pesticides monitored for less than two years are indicated by an asterisk and pesticides monitored for less than one year are not included.

below drainage depth. Comparably high levels of leaching of tebuconazole have not been observed with any of the applications at the three other PLAP sites (Table 17).

The monitoring data also indicate leaching of a further 18 pesticides, but not in similar high concentrations. Although the concentrations exceeded 0.1 µg/l in several samples, the average leaching concentration (1 m b.g.s.) did not. This is summarized in Table 18, showing the number of samples in which the various pesticides were detected on each site as well as the maximum concentration. Apart from slight leaching of ETU (Kjær *et al.*, 2002) and amidosulfuron, leaching within this group of pesticides was only observed at the loamy soil sites, where it was associated with pronounced macropore transport, resulting in very rapid movement of pesticides through the unsaturated zone. It should be noted that the findings regarding amidosulfuron are of very limited use since the degradation product – with which the leaching risk is mainly associated – are not included, as methods for their analysis are not yet available.

Ten of the 40 pesticides applied – about 25% – did not leach during the monitoring period. This group includes the three different sulfonylureas – metsulfuron-methyl, triasulfuron, iodosulfuron-methyl-sodium and tribenuron-methyl applied at several sites. For example, tribenuron-methyl was applied at four different sites under different hydrological conditions, percolation (1 m b.g.s.) during the first month after application ranging from 0 to 114 mm. The monitoring results give no indication of leaching for

any of the compounds or their degradation products. It should, however, be noted, that the leaching risk associated with an autumn application of tribenuron-methyl, where preferential transport is likely to occur, has not yet been evaluated for the loamy soils.

The leaching patterns of the sandy and loamy sites are further illustrated in Figure 40 and 41, showing the frequency of detection in samples collected 1 m b.g.s. (suction cups on sandy soils and drainage systems on loamy soils) and the deeper located groundwater monitoring screens.

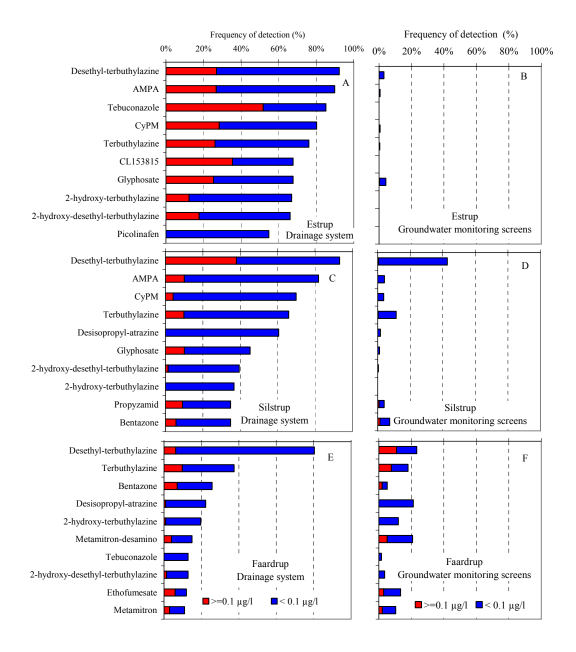


Figure 41. Frequency of detection in samples from the drainage system (left) and groundwater monitoring screens located deeper than the drainage system (right) at the loamy soil sites Silstrup (A, B), Estrup (C, D) and Faardrup (E, F). Frequency is estimated for the entire monitoring period and the time that the different pesticides have been included in the programme and the number of analysed samples thus varies considerably among the different pesticides. The figure only includes the ten most frequently detected pesticides. Pesticides monitored for less than two years are indicated by an asterisk and pesticides monitored for less than one year are not included.

On the sandy soils, as compared to the loamy, the number of leached pesticides as well as the frequency of detection was much lower (Figure 40 and 41), the exceptions being the mobile and persistent degradation products of rimsulfuron and metribuzin, frequently found in both suction cups and groundwater monitoring wells. This difference was mainly due to the different flow patterns characterising the two different soil types. On the sandy soils infiltrating water mainly passed through the matrix, thereby providing good conditions for sorption and degradation. Pesticides being leached in the sandy soils were thus restricted to mobile as well as persistent pesticides. On the loamy soils pronounced macropore transport resulted in the pesticides moving very rapidly through the unsaturated zone. Compared to the sandy soils residence time was much lower on the structured, loamy soils. As a result of this, various types of pesticides, even those being strongly sorbed, were prone to leaching on the loamy types of soil.

At the loamy sites pronounced leaching was generally confined to the depth of the drainage system. Several pesticides were often detected in the drainage system, whereas the amount of pesticide reaching the monitoring screens situated beneath the drainage system was limited and varied considerably within the three sites (Figure 41). These differences should be seen in relation to the different sampling procedures applied. Frequent, integrated water samples can be provided from a drainage system that continuously capture water infiltrating throughout the drainage runoff season. However, although the monitoring screens situated beneath the drainage systems were sampled less frequently (on a monthly basis from a limited number of the monitoring screens (Appendix 2), pesticides were frequently found in selected screens at Faardrup and Silstrup. Hitherto, at the Estrup site leaching of pesticides has mainly been confined to the depth of the drainage system. Apart from 25 samples containing glyphosate, pesticides have only sporadically been detected in the screens beneath the drainage system (Appendix 5). The differences are, however, largely attributable to the hydrological conditions. Compared to the Silstrup and Faardrup sites, the C horizon (situated beneath the drainage depth) at the Estrup site is less permeable with less preferential flow through macropores (se Kjær et al. 2005c for details). The movement of water and solute may therefore be slower at Estrup, allowing for dispersion, dilution, sorption and degradation and thereby reducing risk of a deeper transport.

Comparing the loamy sites, the number of drainage water samples containing pesticides/degradation products was markedly higher at Silstrup and Estrup than at Faardrup, largely attributable to the differences in the hydrological conditions; precipitation and subsequent percolation occurring within the first month after application were, generally, higher at Silstrup and Estrup than at Faardrup (Table 9, Table 11 and Table 13).

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Table A1.1 Systematic chemical nomenclature for the pesticides and degradation products encompassed by the PLAP

PLAP.	
Parameter	Systematic chemical nomenclature
AIBA*	2-amino-N-isopropyl-benzamid
AMPA	Amino-methylphosphonic acid
Amidosulfuron	N-[[[(4,6-dimethoxy-2-pyrimidinyl)amino]carbonyl]-amino]sulfonyl]-N-methylmethanesulfonamide
Azoxystrobin	Methyl (E)-2-{2-[(6-(2-cyanophenoxy)-4-pyrimidin-4-yloxy]phenyl}-3-methoxyacrylate
Bentazone	3-(1-methylethyl)-1H-2,1,3-benzothiadiazin-4(3H)-one 2,2 dioxide
Bromoxynil	3,5-dibromo-4-hydroxybenzonitrile
CGA 322704 *	N-(2-chloro-thiazol-5-yl-methyl)-N'-methyl-N"-nitro-guanidine
Chlormequat	2-chloroethyltrimethylammonium chloride
Clomazone	2-[(2-chlorphenyl)methyl]-4,4-dimethyl-3-isoxazolidione
CL153815	6-(3-trifluoromethylphenoxy)-2-pyridine carboxylic acid
Clopyralid	3,6-Dichloropyridine-2-carboxylic acid
CyPM*	E-2-(2-[6-cyanophenoxy)-pyrimidin-4-yloxy]-phenyl) – 3-methoxyacrylic acid
Desethyl-terbuthylazine *	6-chloro-N-(1,1-dimethylethyl)-1,3,5,triazine-2,4-diamine
Desisopropyl-atrazine *	6-chloro-N-ethyl-1,3,5,triazine-2,4-diamine
Desmedipham	Ethyl 3-(phenylcarbamoyloxy)phenylcarbamate
Dimethoate	O,O-dimethyl S-methylcarbamoylmethyl-phosphorodithioate
EHPC *	Ethyl 3-hydroxy-phenylcarbamate
Epoxiconazole*	(2RS, 3SR)-1-(2-(2-chlorophenyl)-2,3-epoxy-2-(4-fluorophenyl)propyl)-1H-1,2,4-triazol
Ethofumesate	(±)-2-ethoxy-2,3-dihydro-3,3-dimethylbenzofuran-5-yl-methanesulfonate
ETU *	Ethylenethiourea
Fenpropimorph	Cis-4-[3-[4-(1,1-dimethylethyl)-phenyl]-2-methylpropyl]-2,6-
	dimethylmorpholine
Fenpropimorphic acid*	Cis-4-[3-[4-(2-carboxypropyl)-phenyl]-2-methylpropyl]-2,6-
TI (C :1)*	dimethylmorpholine
Flamprop (free acid)*	N-benzoyl-N-(3-chloro-4-flourophenyl)-D-alanine
Flamprop-M-isopropyl	Isopropyl N-benzoyl-N-(3-chloro-4-flourophenyl)-D-alaninate 2',6',8-Trifluoro-5-methoxy-s-triazolo [1,5-c]pyrimidine-2-sulfonanilide
Florasulam	2 ,6 ,8-11111do10-3-methoxy-s-triazoto [1,5-c]pyrimidine-2-surionaniide
Florasulam-desmethyl *	N-(2,6-difluorophenyl)-8-fluro-5-hydroxy[1,2,4]triazolo[1,5-c]pyrimidine-2-sulfonamide
Fluazifop-P-butyl	Butyl (R)-2-[4-(5-trifuoromethyl-2-pyridyloxy)phenoxy]propionate
Fluazifop-P (free acid)*	(R)-2-[4-(5-trifluoromethyl-2-pyridyloxy)phenoxy]propanoic acid
Fluroxypyr	(4-amino-3,5-dichloro-6-fluro-2-pyridinyl)oxy]acetic acid
Glyphosate	N-(phosphonomethyl)glycine
2-hydroxy-terbuthylazine*	6-hydroxy-N-(1,1-dimethylethyl)-N'-ethyl-1,3,5,triazine-2,4-diamine
2-hydroxy-desethyl-terbuthylazine*	6-hydroxy-N-(1,1-dimethylethyl)-1,3,5,triazine-2,4-diamine
Ioxynil	4-hydroxy-3,5-diiodobenzonitrile
Iodosulfuron-methyl	methyl-4-iodo-2-[3-(4-methoxy-6-methyl-1,3,5-triazin-2-yl)-
	ureidosulfonyl]benzoate
Linuron	3-(3,4-dichlorophenyl)-1-methoxy-1-methylurea
MCPA	(4-cloro-2-methylphenoxy)acetic acid
Mesosulfuron	$2-[(4,6-dimethoxypyrimidin-2-ylcarbamoyl)sulfamoyl]-\alpha-$ (methanesulfonamido)-p-toluic acid
Mesosulfuronmethyl *	Methyl 2-[3-(4,6-dimethoxypyrimidin-2-yl)ureidosulfonyl]-4-
Metamitron	methanesulfonamidomethylbenzoate 4-amino-4,5-dihydro-3-methyl-6-phenyl-1,2,4-triazin-5-one
Metamitron-desamino*	4,5-dihydro-3-methyl-6-phenyl-1,2,4-triazine-5-one
Metribuzine	4,5-dinydro-5-methyl-o-pnenyl-1,2,4-triazine-5-one 4-amino-6-tert-butyl-4,5-dihydro-3-methylthio-1,2,4-triazine-5-one
Metribuzine-desamino-diketo*	6-tert-butyl-4,5-dihydro-3-methylthio-1,2,4-triazine-3,5-dione
* Degradation product	o terrossij. 1,5 dinjuro 5 mempuno 1,2,1 unizino 5,5 dione

^{*} Degradation product

Table A1.1 (continued) Systematic chemical nomenclature for the pesticides and degradation products encompassed by the PLAP.

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Parameter	Systematic chemical nomenclature
Metribuzine-desamino	6-(1,1-dimethylethyl)-3-(methylthio)-1,2,4-triazin-5(4H)-one
Metribuzine-diketo*	4-amino-6-tert-butyl-4,5-dihydro-1,2,4-triazine-3,5-dione
Metsulfuron-methyl	Methyl2-[[[(4-methoxy-6-methyl-1,3,5-triazine-2-yl)amino]=carbonyl]amino]-
	sulfonyl]benzoic acid
MHPC*	Methyl-N-(3-hydoxyphenyl)-carbamate
Pendimethalin	N-(1-ethyl)-2,6-dinitro-3,4-xynile
Phenmedipham	3-[(methoxycarbonyl)amino]phenyl (3-methylphenyl)carbamate
PHCP*	3-phenyl-4-hydroxy-6-chloropyridazine
Picolinafen	4'-fluoro-6-(α,α,α-trifluoro-m-tolyloxy)pyridine-2-carboxanilide
Pirimicarb	2-(dimethylamino)-5,6-dimethyl-4-pyrimidinyldimethylcarbamate
Pirimicarb-desmethyl*	2-(dimethylamino)-5,6-dimethyl-4-pyrimidinylmethylcarbamate
Pirimicarb-desmethyl-	2-methylformamido-5,6-dimethylpyrimidine-4-yl dimethylcarbamate
formamido*	
PPU*	N-(4,6-dimethoxy-2-pyrimidinyl-N-((3-ethylsulfonyl)-2-pyridinyl)urea (IN70941)
PPU-desamido*	N-((3-(ethylsulfonyl)-2-pyridyl)-4,6 dimethoxy-2 pyrimidinamine (IN70942)
Propanamide-clomazone*	(N-[2- chlorophenol)methyl] -3-hydroxy-2,2- dimethyl propanamide
Propiconazole	1-[[2-(2,4-dichlorophenyl)-4-propyl-1,3-dioxolan-2-yl]methyl]-1H-1,2,4-triazole
Propyzamide	3,5-dichloro-N-(1,1-dimethylprop-2-ynyl)benzamide
Prosulfocarb	N-[[3-(4-methoxy-6-methyl-1,3,5-triazin-2-yl)-3-[2-(3,3,3,-
	trifluro=propyl)phenylsulfonyl]urea
Rimsulfuron	N-[[(4,6-dimethoxy-2-pyrimidinyl)amino]carbonyl]-3-(ethylsulfonyl)-2-
	pyridinesulfonamide
RH-24580*	N-(1,1-dimethylacetonyl)-3,5-dichlorobenzamide
RH-24644*	2-(3,5-dichlorophenyl)-4,4-dimethyl-5-methylene-oxalzoline
RH-24655*	3,5-dichloro-N-(1,1-dimethylpropenyl)benzamide
Terbuthylazine	6-chloro-N-(1,1-dimethylethyl)-N-ethyl-1,3,5,triazine-2,4-diamine
Tebuconazole	a-[2-(4-chlorophenyl)ethyl]-a-(1,1-dimethylethyl)-1H-1,2,4-triazole-1-ethanol
Thiamethoxam	3-(2-cholro-thiazol-5-ylmethyl)-5-methyl[1,3,5]oxadiazinan-4ylidene-N-nitroamine
Triasulfuron	1-[2-(2-chloroethoxy)phenylsulfonyl]-2-(4-methoxy-6-methyl-1,3,5-triazine-2-yl)-urea
Triazinamin	4-methoxy-6-methyl-1,3,5-triazine-2-amin
Triazinamin-methyl*	1,3,5-triazine-2-2-amine 4-methoxy-N, 6-dimethyl
i D	130,0 and 2.2 and 1 memory 11, 0 and my

^{*} Degradation product

From each of the PLAP sites, samples were collected of groundwater, drainage water and soil water in the unsaturated zone. A full description of the monitoring design and sampling procedure is provided in Lindhardt *et al.* (2001) and Kjær *et al.* 2003 respectively.

Until March 2002, pesticide analysis was performed monthly on water samples from the suction cups located both 1 m b.g.s. and 2 m b.g.s., from two screens of the horizontal monitoring wells and from two of the downstream vertical monitoring wells. In addition, more intensive monitoring encompassing all four groups of suction cups, six screens of the horizontal monitoring wells and five monitoring wells was performed every fourth months (Kjær *et al.*, 2002). At the loamy sites, the pesticide analysis was also performed on drainage water samples. The monitoring programme was revised in March 2002 and the number of pesticide analyses was reduced. At the loamy sites, pesticide analysis of water sampled from the suction cups was ceased, and the monthly monitoring was restricted to just one monitoring well. At Jyndevad, pesticide analysis of the suction cups located 2 m b.g.s. was ceased and the interval for the intensive monitoring encompassing the larger number of monitoring screens was extended to six months, except for the suction cups 2 m b.g.s. at Tylstrup, where the four-month interval was retained (Kjær *et al.*, 2003).

On the sandy soils, the analysis of a number of pesticides in the monitoring wells had to be further reduced, due to economical constraints imposed by the high prices on pesticide analysis (see Table A5.1 and Table A5.2 in Appendix 5). However, results from the suction cups implied that the leaching risk of those pesticides was negligible, and followingly they were only measured in a limited number of groundwater samples.

Until July 2004, pesticide analyses were performed weekly on water sampled time-proportionally from the drainage system. Moreover, during storm events additional samples (sampled flow-proportionally during 1-2 days) were also analysed for pesticides. In June 2004 the drainage monitoring programme was revised. From July 2004 and onwards pesticide analyses were done weekly on water sampled flow-proportionally from the drainage water system. See Kjær *et al.* 2003 for further details on the methods of flow-proportional sampling. The weighted average concentration of pesticides in the drainage water was calculated according to the following equation:

$$C = \frac{\sum_{i=1}^{n} M_i}{\sum_{i=1}^{n} V_i}$$

$$M_i = C_i \cdot V_i$$

where:

n = Number of weeks within the period of continuous drainage runoff

 V_i = Weekly accumulated drainage runoff (mm/week)

 C_i = Pesticide concentration collected by means of the flow-proportional sampler ($\mu g/l$)

Until July 2004 where both time and flow-proportional sampling was applied the numbers were:

 $M_i = Ct_i \cdot V_i$ If no flow event occurs within the i'th week $M_i = Cf_i \cdot Vf_i$ If a flow event occurs within the i'th week and if $Cf_i \cdot Vf_i > Ct_i \cdot V_i$

where:

n = Number of weeks within the period of continuous drainage runoff

V_i = Weekly accumulated drainage runoff (mm/week)

Vf_i = Drainage runoff accumulated during a "flow event" (mm/storm event)

 Cf_i = Pesticide concentration in the "event samples" collected by means of the flow-proportional sampler ($\mu g/l$)

 Ct_i = Pesticide concentration in the weekly samples collected by means of the time-proportional sampler ($\mu g/l$)

Table 9, 10 and 11 report the weighted average leachate concentration in the drainage water within the first drainage season after application. In these tables this calculation period is defined as the period from the date of application until 1 July the following year.

On the sandy soils the weighted average concentration of pesticides leached to the suction cups situated 1 m b.g.s. was estimated using the measured pesticide concentration and estimated percolation on a monthly basis. Pesticide concentrations measured in suction cups S1 and S2 were assumed to be representative for each sample period. Moreover, accumulated percolation rates deriving from the MACRO model were assumed to be representative for both suction cup S1 and suction cup S2. For each of the measured concentrations, the corresponding percolation (Perc.) was estimated according to the equation:

$$P_i = \sum_{t_1}^{t_2} P_t$$

where

 $t = \text{sampling date}; t_1 = 0.5(t_{i-1}+t_i); t_2=0.5(t_i+t_{i+1})$

 P_t = daily percolation at 1 m b.g.s. as estimated by the MACRO model (mm)

The average concentration was estimated according to the equation:

$$C = \frac{\sum C_i \cdot P_i}{\sum P_i}$$

where

 C_i = measured pesticide concentration in the suction cups located 1 m b.g.s.

Table A3.1 Management practice at Tylstrup during the 2005 to 2008 growing seasons. The active ingredients of the various pesticides are indicated in parentheses.

Date	Management practice
10.10.05	Maize harvested (harvest yield 137.75 hkg/ha, 100% DM)
05.11.05	Rotary cultivated - 8 cm depth
21.04.06	Ploughed - 22 cm depth
21.04.06	Rolled with a concrete roller
23.04.06	Seed bed preparation – 7 cm depth
23.04.06	Spring barley sown (cultivare Cabaret)
06.06.06	Herbicide - 4.25 tablets/ha Express ST (tribenuron-methyl)
03.07.06	Fungicide - Opus (epoxiconazole) - 1.0 L/ha
19.06.05	Irrigation - 29 mm
24.08.06	Spring barley harvested (grain yield 55,6 hkg/ha 85% DM)
24.08.06	Straw shredded, amount 33.2 hkg/ha, 100% DM
25.08.06	Rotary cultivated - depth 5.0 cm (straw incorporation)
26.08.06	Ploughed - depth 23 cm.
26.08.06	Winter rape sown (cv. Lioness)
26.08.06	Herbicide - 0.33 L/ha Command CS (clomazone)
06.09.06	Seedbed preparation – 7 cm depth
06.09.06	Winter rape resown (cv. Castille)
09.02.07	Herbicide - 1.0 l/ha Kerb 500 SC (propyzamide)
27.03.07	Herbicide - 0.8 ll/ha Matrigon (clopyralid)
08.06.07	Irrigation 30 mm
01.00.07	Direct harvest and simultaneous shredding of straw (Seed yield 24.5 hkg/ha 91% DM, straw
01.08.07	amount 64.7 hkg/ha 100% DM incl. stubble)
03.08.07	Rotary cultivated - depth 3.0 cm (straw incorporation)
07.09.07	Rotary cultivated - depth 7.0 cm (straw incorporation)
12.09.07	Ploughed - 22 cm depth
12.09.07	Winter wheat sown, cv. Smuggler.
18.10.07	Herbicide - 5.0 l/ha Stomp (pendimethalin)
16.11.07	Herbicide - 1.0 l/ha Folicur EC250 (tebuconazole)
22.05.08	Irrigation 32 mm
29.05.08	Irrigation 32 mm
05.06.08	Irrigation 32 mm
13.06.08	Irrigation 30 mm
17.06.08	Fungicide – 1.0 l/ha Amistar (azoxystrobin)

Table A3.2 Management practice at Jyndevad during the 2005 to 2008 growing seasons. The active ingredients of the various pesticides are indicated in parentheses.

Date	Management practice
19.08.05	Winter wheat harvested (grain yield 57.5 hkg/ha, 85% DM. Straw yield 34.0 hkg/ha, 100% DM)
29.03.06	Rotary cultivated - 5 cm depth
30.03.06	Ploughed - 20 cm depth
04.04.06	Rolled with a concrete roller
11.04.06	Spring barley sown - cv. Simba
26.05.06	Herbicide - 0.1 l/ha Primus (florasulam)
06.06.06	Irrigation - 27 mm
08.06.06	Herbicide – 1.0 l/ha Opus (epoxiconazole)
12.06.06	Irrigation - 30 mm
26.06.06	Irrigation - 27 mm
02.07.06	Irrigation - 30 mm
07.07.06	Irrigation - 30 mm
07.08.06	Spring barley harvested (Seed yield 60.49 kgh/ha; 85% DM. Straw yield 26.75 hkg/ha 100% DM)
20.09.06	Ploughed - 22 cm depth
20.09.06	Rolled with a concrete roller
21.09.06	Seed bed preparation - 6 cm depth
21.09.06	Triticale sown - cv. Dinaro
10.10.06	Herbicide - 250 g/ha Atlantis WG (mesosulfuron-methyl/iodosulfuron)
13.04.07	Plant growth inhibitor – 1.0 l/ha Cycocel 750 (chlormequat)
27.04.07	Irrigation - 27 mm
07.05.07	Herbicide - 1.0 l/ha Opus (epoxiconazole)
05.06.07	Irrigation - 27 mm
07.08.07	Harvest of triticale (seed yield 38.7 kgh/ha 85% DM, straw yield 38.3 hkg/ha 100% DM)
13.09.07	Herbicide - 2.0 l/ha Roundup (glyphosate, not monitored)
28.09.07	Ploughed - 22 cm depth
29.09.07	Rolled with a concrete roller
01.10.07	Winter wheat sown – cv. Ambition
29.10.07	Herbicide - 0.133 g/ha Pico 750 WG (picolinafen)
03.12.07	Fungicide - 1.0 l/ha Folicur EC 250 (tebuconazole)
07.05.08	Irrigation - 42 mm
14.05.08	Irrigation - 27 mm
21.05.08	Irrigation - 27 mm
30.05.08	Irrigation - 30 mm
05.06.08	Irrigation - 35 mm
11.06.08	Fungicide - 1.0 l/ha Amistar (azoxystrobin)
25.06.08	Irrigation - 35 mm

Table A3.3 Management practice at Silstrup during the 2005 to 2008 growing seasons. The active ingredients of the various pesticides are indicated in parentheses.

Date	Management practice
01.09.05	Winter rape sown – cultivare Calypso
03.09.05	Herbicide – 0.33 l/ha Command CS (Clomazone)
17.11.05	Herbicide – 1.0 l/ha Kerb 500 SC (Propyzamide)
17.11.05	Herbicide Kerb 500 SC (propyzamide - 1.0 l/ha)
20.04.06	Herbicide –0.8 l/ha Matrigon (clopyralid)
05.05.06	Insecticide - 0.25 l/ha Fastac (alpha-cypermethin, not analyzed)
24.07.06	Windrowed – stubleheight 13 cm
07.08.06	Threshed (seed yield 37.3 hkg/ha 91% DM)
20.09.06	Ploughed - 22 cm depth
22.09.06	Winter wheat sown - cv. Skalmeje
22.09.06	Herbicide - 5.0 l/ha Stomp Pentagon (pendimethalin)
13.04.07	Herbicide - 100 ml/ha Husar OD (iodosulfuron)
13.04.07	Growth retardent - 1.2 L/ha Cycocel 750 (chlormequat)
07.06.07	Fungicide - 1.0 l/ha Opus (epoxiconazole)
24.08.07	Winter wheat harvested. Grain yield 100.7 hkg/ha 85% DM, straw yield 40.8 hkg/ha 100% DM, shredded at harvest
29.08.07	Stubble harrowed, heavy disk harrow (Dalbo) - 5 cm depth
12.11.07	Ploughed - 27 cm depth
21.04.08	Harrowed - depth 5 cm and packed
05.05.08	Harrowed across - depth 10 cm
07.05.08	Fodderbeet sown - cv. Kyros.
22.05.08	Herbicide - 30 g/ha Safari (triflusulfuron-methyl) + 0.5 l/ha Goliath (metamitron) + 1.5 l/ha Betanal (phenmedipham)
30.05.08	Herbicide - 30 g/ha Safari (triflusulfuron-methyl) + 0.5 l/ha Goliath (metamitron) + 1.5 l/ha Betanal (phenmedipham) + 0.07 l/haTramat 500 SC (ethofumesat)
17.06.08	Herbicide - 30 g/ha Safari (triflusulfuron-methyl) + 0.5 l/ha Goliath (metamitron) + 1.5 l/ha Betanal (phenmedipham) + 0.07 l/haTramat 500 SC (ethofumesat)
26.06.08	Insecticide - 0.30kg/ha Pirimor G (pirimicarb)

Table A3.4 Management practice at Estrup during the 2005 to 2008 growing seasons. The active ingredients of the various pesticides are indicated in parentheses.

1	cides are indicated in parentheses.	
Date	Management practice	
13.10.05	Maize harvested (harvest yield 143.4 hkg/ha, 100% dry matter)	
12.04.06	Ploughed – depth 18 cm - packed with a ring roller	
07.04.06	Spring barley sown – cv. Simba	
01.05.06	Rolled with a cam roller	
17.05.06	Herbicide – 0.25 l/ha Starane 180 (fluroxypyr, not analyzed)	
02.06.06	Herbicide – 0.25 l/ha Starane 180 (fluroxypyr, not analyzed)	
06.06.06	Herbicide – 0.10 l/ha Primus (florasulam)	
29.06.06	Fungicide - 1.0 l/ha Amistar (azoxystrobin)	
17.08.06	Harvest of spring barley (grain yield 59.2 hkg/ha; 85% DM)	
11.09.06	Straw removed (straw yield 26.79 hkg/ha, 100% DM)	
13.09.06	Ploughed - 18 cm depth (packed with a ring roller)	
14.09.06	Rotary cultivated - 4 cm depth	
14.09.06	Winter wheat sown – cv. Smuggler	
11.10.06	Herbicide - 250 g/ha Atlantis WG (mesosulfuron/iodosulfuron)	
11.04.07	Growth retardent - 1.2 l/ha Cycocel 750 (chlormequat)	
31.05.07	Fungicide - 1.0 l/ha Opus (epoxiconazole)	
07.08.07	Winter wheat harvested (Grain yield 81.5 hkg/ha, 85% DM)	
08.08.07	Straw shredded (47.4 hkg/ha, 100 % DM)	
02.10.07	Ploughed - depth 20 cm (packed with a ring roller)	
02.10.07	Seed bed preparation (rotary harrowed across, depth 4 cm)	
03.10.07	Winter wheat sown – cv. Frument.	
30.10.07	Herbicide - 0.133 g/ha Pico 750 WG (picolinafen)	
22.11.07	Fungicide - 1.0 l/ha Folicur EC 250 (tebuconazole)	
13.06.08	Fungicide - 1.0 Amistar (azoxystrobin)	

Table A3.5 Management practice at Faardrup during the 2005 to 2008 growing seasons. The active ingredients of the various pesticides are indicated in parentheses.

Date	Management practice
28.09.05	Maize harvested (harvest yield 160.3 hkg/ha, 100% DM)
30.11.05	Ploughed – 25 cm depth
28.04.06	Spring barley sown - cv. Scandium
26.05.06	Herbicide - 0.8 l/ha Starane 180 S (fluroxypyr)
29.06.06	Fungicide - 1.0 l/ha Opus (epoxiconazole)
11.08.06	Harvest of spring barley (grain yield 67.3 hkg; 85% DM. Straw yield 51.1 hkg/ha 100% DM)
17.08.06	Ploughed - 22 cm depth
17.08.06	Winter rape sown – cv. Labrador
18.08.06	Herbicide - 0.33 l/ha Command CS (clomazone – not analysed)
19.02.07	Herbicide - 1.0 l/ha Kerb 500 SC (propyzamide)
30.03.07	Herbicide – 0.8 l/ha Matrigon (clopyralid)
03-07-07	Windrowed, stubble height 20 cm
19.07.07	Threshed winter rape. Seed yield 30.0 hkg/ha 91% DM
19.07.07	Straw shredded. 68.5 hkg/ha 100% DM
10.08.07	Stubble cultivation - 15 cm depth
22.08.07	Stubble cultivation - 15 cm depth
18.09.07	Ploughed and packed - depth 25 cm.
18.09.07	Winter wheat sown – cv. Ambition
09.10.07	Herbicide - 5.0 Stomp (pendimethalin)
20.11.07	Fungicide - 1.0 L/ha Folicur 250 (tebuconazole)

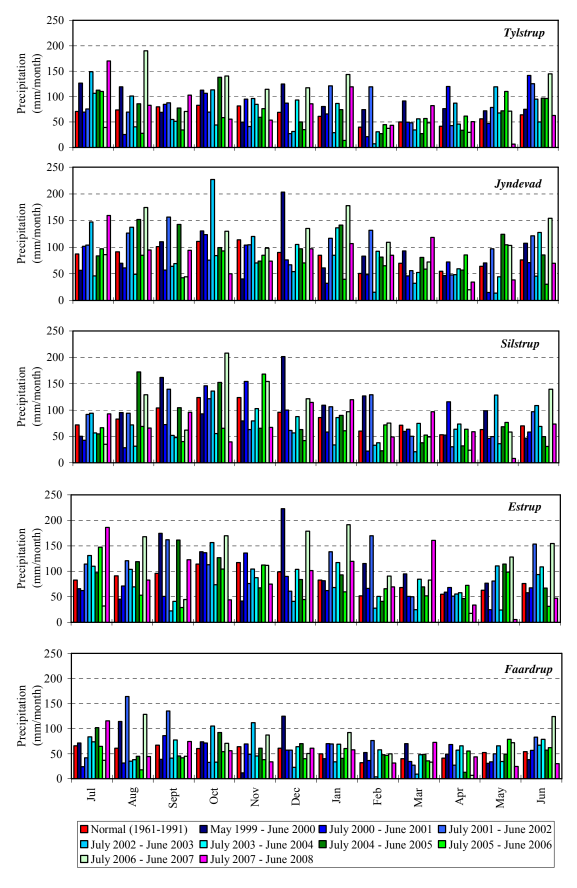


Figure A4.1. Monthly precipitation at all localities for the monitoring period July 2000–June 2008. Normal values (1961–1990) are included for comparison.

Table A5.1 Number of samples where pesticides were either not detected (n.d.), detected in concentrations below 0.1 μ g/l (det<0.1 μ g/l) or detected in concentrations above 0.1 μ g/l (det>=0.1 μ g/l) at Tylstrup. Numbers are accumulated for the entire monitoring period, and pesticides monitored for less than one year are not included.

		Vertical so	reens	Suction cups						
	n.d.	$det.{<}0.1~\mu g/l$	det.>=0.1 μ g/l	n.d.	$det.{<}0.1~\mu g/l$	det.>=0.1 μ g/l				
2-amino-N-isopropylbenzamid	191			72						
2-hydroxy-desethyl-terbuthylazine	190	1		67	5					
2-hydroxy-terbuthylazin	191			71	1					
Bentazone	191			71	1					
Bromoxynil	192			72						
CGA 322704	175			64						
Clomazone	224			82						
Clopyralid*	6			40						
CyPM	13			6						
Desethyl-terbuthylazine	191			70	2					
Desisopropyl-atrazine	190	1		55	17					
Dimethoate	176			65						
Epoxiconazole	199			74						
ETU	198	2		37	7					
Fenpropimorph	307			89						
Fenpropimorph-acid	276			73						
Flamprop (free acid)	176			65						
Flamprop-M-isopropyl	176			65						
Fluazifop-P (free acid)	178			65						
Fluroxypyr	194			70						
FMC65317	208			74						
IN70941	358	1		63	65	3				
IN70942	359			102	29					
Ioxynil	198			72						
Linuron	270			67						
Metribuzine	386	1		89	2					
Metribuzine-desamino	365			85						
Metribuzine-desamino-diketo	289	231	5	168	30	51				
Metribuzine-diketo	71	136	317	44	147	63				
Pendimethalin	307			96						
Pirimicarb	295			82						
Pirimicarb-desmethyl	295			81						
Pirimicarb-desmethyl-formamido	167			52						
Propiconazole	307			89						
Propyzamide	157			58						
RH24580	157			58						
RH24644	157			58						
RH24655	157			58						
Rimsulfuron	178			65						
Tebuconazole	58	1		24						
Terbuthylazine	179			72						
Thiamethoxam	175			64						
Triasulfuron	295			82						
Triazinamin	285			75						
Triazinamin-methyl	440			137						

^{*}Number of analysed samples collected from the monitoring wells was reduced (see Appendix 2 for explanation).

Table A5.2 Number of samples where pesticides were either not detected (n.d), detected in concentrations below 0.1 μ g/l (det<0.1 μ g/l) or detected in concentrations above 0.1 μ g/l (det>=0.1 μ g/l) at Jyndevad. Numbers are accumulated for the entire monitoring period, and pesticides monitored for less than one year are not included.

		Vertical so	reens	Suction cups					
	n.d.	$det.{<}0.1~\mu g/l$	det.>=0.1 μ g/l	n.d.	$det.{<}0.1~\mu g/l$	det.>=0.1 μ g/1			
2-amino-N-isopropylbenzamid	178			45	2				
4-chlor-2-methylphenol	189			52					
Amidosulfuron	88			20	2	1			
AMPA	221	2		68	1				
Azoxystrobin	233			65					
Bentazone	277			47	27	3			
Bromoxynil	218			61					
CL153815*				14					
Chlormequat	14			28					
CyPM	233			65					
Desethyl-terbuthylazine	472	24		128	18				
Desmethyl-amidosulfuron	88			23					
Dimethoate	169			48					
Epoxiconazole	192			54					
Fenpropimorph	246	1		76	1				
Fenpropimorph-acid	259			79					
Flamprop (free acid)	12			4					
Flamprop-M-isopropyl	12			4					
Florasulam	192			54					
Florasulam-desmethyl*				28					
Fluazifop-P (free acid)	190			51					
Fluroxypyr	193			55					
Glyphosate	223			69					
IN70941	363	155	2	6	87	50			
IN70942	470	50		56	83	4			
Ioxynil	218			61					
MCPA	189			52					
Mesosulfuron*				25					
Mesosulfuron-methyl	154			42					
Metribuzine	26			6					
Metribuzin-desamino	26			4					
Metribuzin-desamino-diketo	6	7	13	6					
Metribuzin-diketo		7	19	3	3				
Pendimethalin	257			71					
PHCP	184			59					
Picolinafen*				14					
Pirimicarb	251			69					
Pirimicarb-desmethyl	251			68	1				
Pirimicarb-desmethyl-formamido	251			69					
Propiconazole	236			75					
Pyridate	116			39					
Rimsulfuron	168			48					
Tebuconazole	59			16					
Terbuthylazine	239			75					
Triazinamin-methyl	247			77					

^{*} Number of analysed samples collected from the monitoring wells was reduced (see Appendix 2 for explanation).

Table A5.3 Number of samples where pesticides were either not detected (n.d), detected in concentrations below 0.1 μ g/l (det<0.1 μ g/l) or detected in concentrations above 0.1 μ g/l (det>=0.1 μ g/l) at Silstrup. Numbers are accumulated for the entire monitoring periode, and pesticides monitored for less than one year are not included.

for the entire monitoring periode				ontal screens Vertical screens					Suction cups			
			_					det	reens det			_
	n.d.		det >=0.1	n.d.	det <0.1	det >=0.1	n.d.	<0.1	>=0.1	n.d.	det <0.1	det >=0.1
		μg/l	μg/l		μg/l	μg/l		μg/l	μg/l		μg/l	μg/l
2-amino-N-isopropylbenzamid	64			75			131					
2-hydroxy-desethyl-	40	27		0.5			151					
terbuthylazine	43		1	85			151	1				
2-hydroxy-terbuthylazin	45	26		85			152			26		
3-aminophenol	53			70			170			36		
4-chlor-2-methylphenol	51	40-		67	_		124	4.0				
AMPA	27		15	123	5		227	10		8		
Azoxystrobin	44		_	66	_		121					
Bentazone	58		5	95	5	1	169	12	3			
Clopyralid	44			63			118					
Chlormequat	20			36			67					
CyPM	22		3	101	2		187	9				
Desethyl-terbuthylazine	8		44	102	32		113	127	2			
Desisopropyl-atrazine	28	43		85			148	4				
Desmedipham	101			107	1		240			58		
Dimethoate	81		1	73	1		147			27		
EHPC	68			62			118			20		
Epoxiconazole	19			36			69					
Ethofumesate	87	14	1	111	1		244	3		54	3	2
Fenpropimorph	82			74			148			27		
Fenpropimorph-acid	81	1		74			147			27		
Flamprop (free acid)	73	7		74			148			26		
Flamprop-M-isopropyl	70	11	1	73	1		148			27		
fluazifop-P (free acid)	74			81	1		189			56		
Fluroxypyr	50			74			142					
Glyphosate	81	52	15	128			233	4		8		
Iodosulfuron-methyl-natrium	20			36			69					
MCPA	51			67			123					
Metamitron	77	21	4	103	9		230	15	2	40	9	9
Metamitron-desamino	76	23	3	106	3	3	237	9	1	40	15	4
Metsulfuron methyl	20			36			69					
MHPC	100			106			234			55		
Pendimethalin	90	14		122			223					
PHCP	62		4	66	2		109	8	4			
Phenmedipham	101			108			240			59		
Pirimicarb	160	14		210			433	3		59		
Pirimicarb-desmethyl	173			210			436			59		
Pirimicarb-desmethyl-												
formamido	141			160			308			20		
Propiconazole	76	6		74			148			27		
Propyzamid	43	17	6	75	2	1	143	5	1			
Prosulfocarb	69	4	1	79	1		147					
RH24580	64	2		78			149					
RH24644	51	15		77	1		148	1				
RH24655	66			78			149					
Terbuthylazine	31	51	9	108	5		173	30	1			
Triazinamin	18			36			68					
Triazinamin-methyl	82			74			148			27		

Table A5.4 Number of samples where pesticides were either not detected (n.d), detected in concentrations below 0.1 μ g/l (det<0.1 μ g/l) or detected in concentrations above 0.1 μ g/l (det>=0.1 μ g/l) at Estrup. Numbers are accumulated for the entire monitoring periode, and pesticides monitored for less than one year are not included.

for the entire monitoring periode,	Drainage			Horize				ical sc		Suction cups		
	n.d.	-	det	n.d.	det	det	n.d.	det	det	n.d.	Det	det
		< 0.1	>=0.1			>=0.1		< 0.1	>=0.1			>=0.1
		μg/l	μg/l		μg/l	μg/l		μg/l	μg/l		μg/l	μg/l
2-amino-N-isopropylbenzamid	235	1		79	1		271			5		
2-hydroxy-desethyl-	44	63	23	50			180					
terbuthylazine 2-hydroxy-terbuthylazin	43	71		50			180					
4-chlor-2-methylphenol	101	1		34			112					
Amidosulfuron	98	1		34			109					
AMPA	35	219	93	127			461	4		23		
Azoxystrobin	97	54		64			231	4		23		
Bentazone	165	112		92	11		358	1		3	2	2
Bromoxynil	135	1 1 2		41	11		125	1		3		
CL153815	10	10		11			38			3		
Clopyralid	10	10	11	11			36					
Chlormequat	44	1		18			56					
СуРМ	32	84		62	2		231					
Desethyl-terbuthylazine	10	85		43	7		180					
Desisopropyl-atrazine	59	70		49	1		155	25				
Dimethoate	88	70	1	42	1		160	23		23		
Epoxiconazole	35	11	2	19			69			23		
Ethofumesate	91	27		46			158					
Fenpropimorph	82	1		39			152			23		
Fenpropimorph-acid	82			34			125			17		
Flamprop (free acid)	118	13		55			210			23		
Flamprop-M-isopropyl	111	20		55			210			23		
Florasulam	91			35			125					
Florasulam-desmethyl	80			30			100					
Fluroxypyr	87	1	2	34			120	1				
Glyphosate	112	147		124	1		439	22	2	23		
Ioxynil	118	14		41			125			3		
MCPA	91	9		34			111	1				
Mesosulfuron	73			24			83					
Mesosulfuron-methyl	61	13		27			99					
Metamitron	81	27	15	46			158					
Metamitron-desamino	76	38		46			157					
Metsulfuron methyl	130			55			210			22	1	
Picolinafen	14	17		11			38					
Pirimicarb	159	39		67			225	1		6		
Pirimicarb-desmethyl	191			66			223			6		
Pirimicarb-desmethyl-formamido	198	13	13	76			261			5		
Propiconazole	192	22	3	86			311	2		23		
Tebuconazole	4	9		10			36					
Terbuthylazine	31	65	34	50			179	1				
Triazinamin	125			52			197	1		22		

Table A5.5 Number of samples where pesticides were either not detected (n.d), detected in concentrations below 0.1 μ g/l (det<0.1 μ g/l) or detected in concentrations above 0.1 μ g/l (det>=0.1 μ g/l) at Faardrup. Numbers are accumulated for the entire monitoring periode, and pesticides monitored for less than one year are not included.

for the entire monitoring periode, and pesticides monitored for less than one year are not included.											
	Drainage		Horizontal screens				cal scr	eens	Suction cups		
	n.d.	det	det		det	det	n.d.	det	det	n.d.	det det
			>=0.1			>=0.1			>=0.1		<0.1 >=0.1
2	(0	μg/l	μg/l	1	μg/l	μg/l		μg/l	μg/l		μg/l μg/l
2-amino-N-isopropylbenzamid 2-hydroxy-desethyl-	69	1		61			132				
terbuthylazine	61	8	1	60	1		126	6			
2-hydroxy-terbuthylazin	90	21	1	85	4		164	30			
4-chlor-2-methylphenol	144		1	109	-		254				
AMPA	131	9	1	110			282	2		57	5
Azoxystrobin	108		•	92			194	_		0,	C
Bentazone	52	13	5	58	2	1	125	4	3		
Bromoxynil	101	15		81	_	•	226	•	J	73	
CGA 322704	68			58			126			, 5	
Clomazon	84		1	69			166				
СуРМ	104	4	•	92			194				
Desethyl-terbuthylazine	22	83	7	68	21		149	14	31		
Desisopropyl-atrazine	87	24	1	57	32		166	28	31		
Desmedipham	99	27	1	66	32		165	20		29	
Dimethoate	77			58			148			2)	
EHPC	83			52			123			16	
Epoxiconazole	81			66			143			10	
Ethofumesate	88	6	6	66			134	24	7	27	2
Fenpropimorph	70	U	U	58	1		157	24	,	54	2
Fenpropimorph-acid	70			59	1		157			54	
Flamprop (free acid)	76	1		58			148			34	
Flamprop-M-isopropyl	70	1		56			142				
Fluazifop-P (free acid)	91	4	4	66			159	5	1	26	3
Fluazifop-P-buthyl	99	4	4	66			165	3	1	29	3
Fluroxypyr	155		1	128	1		306			55	
FMC65317	84		1	69	1		166			33	
Glyphosate	137	4	1	109	1		282	2		61	1
Ioxynil	99	1		81	1		225	1		73	1
MCPA	143	1	1	109			255	1		13	
Metamitron	89	8	3	66			141	19	5	29	
Metamitron-desamino	85	11	4	66			117	36	12	29	
MHPC	97	11	1	66			163	1	12	29	
Pendimethalin	32	2	1	21			46	1		29	
Phenmedipham	99	2		66			163	2		29	
Pirimicarb	113	7		90			244	2		52	
Pirimicarb-desmethyl	94	6		66			162	3		29	
Pirimicarb-desmethyl-	94	O		00			102	3		29	
formamido	97	3		66			163	2		29	
Propiconazole	147			116			304	1		54	
Propyzamid	62	2	2	50	1		110				
Prosulfocarb	80			61			126				
RH24580	66			51			110				
RH24644	62	4		51			110				
RH24655	65	1		51			110				
Tebuconazole	27	4		19			41	1			
Terbuthylazine	70	31	11	83	5	1	149	24	21		
TFMP	, ,			1	-	-	2				
Thiamethoxam	68			58			126				
Triazinamin-methyl	77			57			147				
	, , ,			31			1 7/				

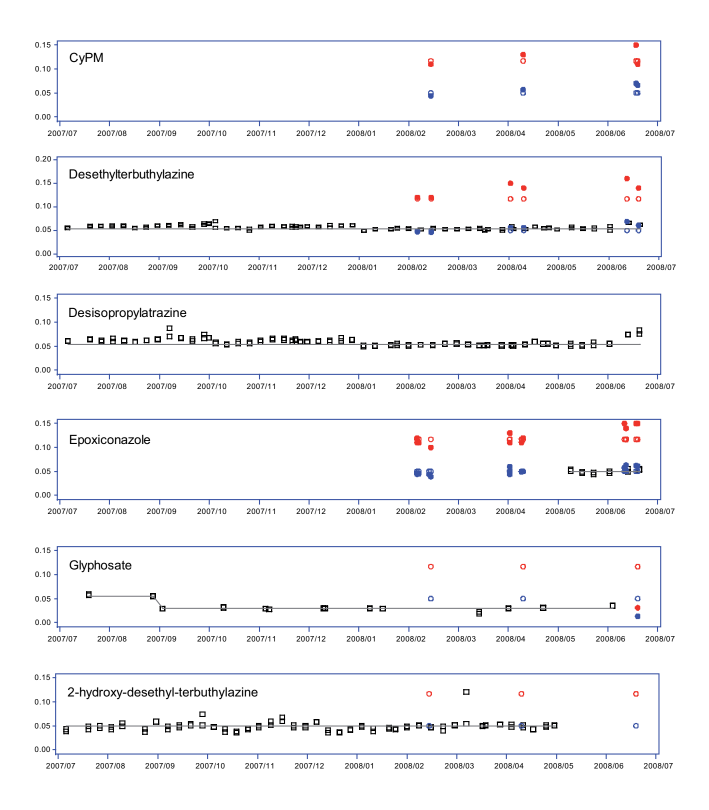


Figure A6.1 Quality control data for pesticide analysis by laboratory 1. <u>Internal laboratory control</u> samples are indicated by square symbols and the nominal level is indicated by the solid grey line (■ IQ measured, — IQ nominal concentration). <u>External control</u> samples are indicated by circles. Open circles indicate the nominal level (○ EQ nominal low, ○ EQ nominal high), and closed circles the observed concentration (● EQ measured low, ● EQ measured high).

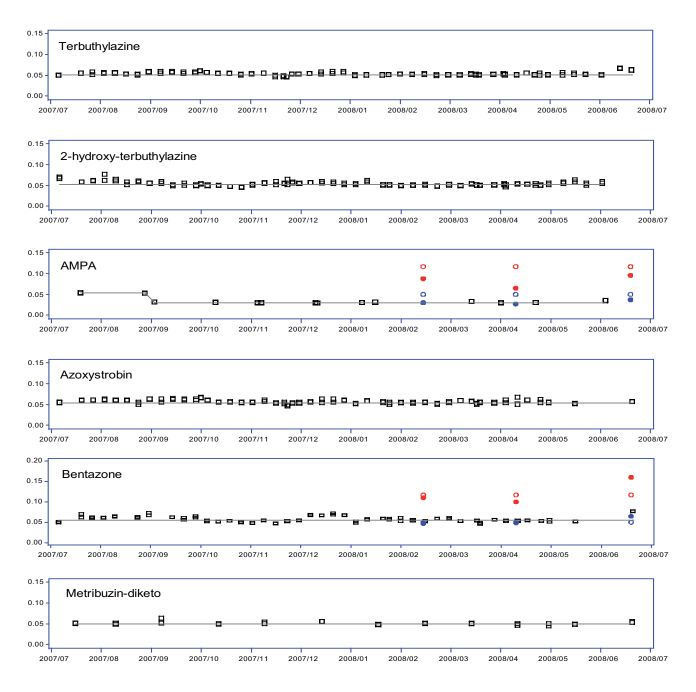


Figure A6.1 (Continued) Quality control data for pesticide analysis by laboratory 1. <u>Internal laboratory control</u> samples are indicated by square symbols and the nominal level is indicated by the solid grey line (■ IQ measured, — IQ nominal concentration). <u>External control</u> samples are indicated by circles. Open circles indicate the nominal level (O EQ nominal low, O EQ nominal high), and closed circles the observed concentration (● EQ measured low, O EQ measured high).

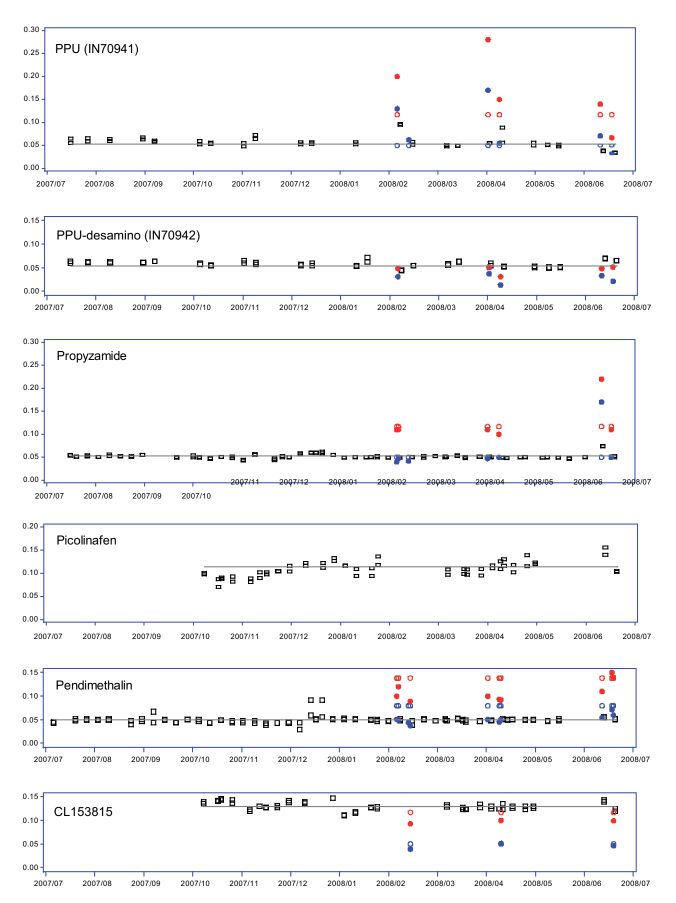


Figure A6.1 (Continued) Quality control data for pesticide analysis by laboratory 1. <u>Internal laboratory control</u> samples are indicated by square symbols and the nominal level is indicated by the solid grey line (■ IQ measured, — IQ nominal concentration). <u>External control</u> samples are indicated by circles. Open circles indicate the nominal level (○ EQ nominal low, ○ EQ nominal high), and closed circles the observed concentration (● EQ measured low, ● EQ measured high).