



# Glucosinolates and their potential role in plant

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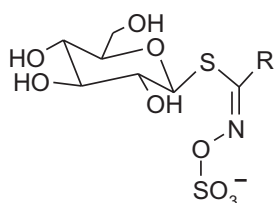
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## Abstract

Glucosinolates are sulfur- and nitrogen-containing plant secondary metabolites common in the Brassicaceae and related plant families. In the plant, they coexist with an endogenous  $\beta$ -thioglucosidase (EC 3.2.3.1) called myrosinase, though glucosinolates are stored in the vacuoles of so-called S-cells and myrosinase in separate but adjacent cells. Upon plant tissue disruption, glucosinolates are released at the damage site and become hydrolyzed by myrosinase. The chemical nature of the hydrolysis products depends on the structure of the glucosinolate side chain, plant species and reaction conditions. Biosynthesis of glucosinolates comprises three phases: (i) amino acid chain elongation, in which additional methylene groups are inserted into the side chain, (ii) conversion of the amino acid moiety to the glucosinolate core structure, (iii) and subsequent side chain modifications. Glucosinolate pattern differs between species and ecotype as well as between and even within individual plants, depending on developmental stage, tissue and photoperiod. A number of environmental conditions such as light plant, nutritional status, fungal infection, wounding and insect damage can alter the glucosinolate pattern significantly. The change of the glucosinolate profile by several environmental factors has brought forward different theories regarding their potential roles in the plant. However, the most accepted theory is that the glucosinolate-myrosinase system is involved in defense against herbivores and pathogens. This review summarized recent progress in glucosinolate biosynthesis, degradation and organization of the myrosinase-glucosinolate system. Furthermore, current knowledge of the potential role of glucosinolates in the plant, especially in plant defense, is discussed.

## INTRODUCTION

Glucosinolates are sulfur- and nitrogen-containing plant secondary metabolites common in the order Capparales, which includes the Brassicaceae family with agriculturally important crops, Brassica vegetables, and the model plant *Arabidopsis thaliana*. Glucosinolates have a common core structure containing a  $\beta$ -D-thioglucose group linked to a sulfonated aldoxime moiety and a variable side chain derived from amino acids (Figure 1). Glucosinolates can be divided into three classes based on the structure of different amino acid precursors: 1. aliphatic glucosinolates derived from methionine, isoleucine, leucine or valine, 2. aromatic glucosinolates derived from phenylalanine or tyrosine, and 3. indole glucosinolates derived from tryptophan. The biosynthesis of glucosinolates comprises three phases: (i) amino acid chain elongation, in which additional methylene groups are inserted into the side chain, (ii) conversion of the amino acid moiety to the glucosinolate core structure, (iii) and subsequent side chain modifications (1). More than 130 glucosinolates have been identified. Their structural diversity arises



**Figure 1.** General structure of glucosinolates. *R* denotes the variable side chain from amino acids.

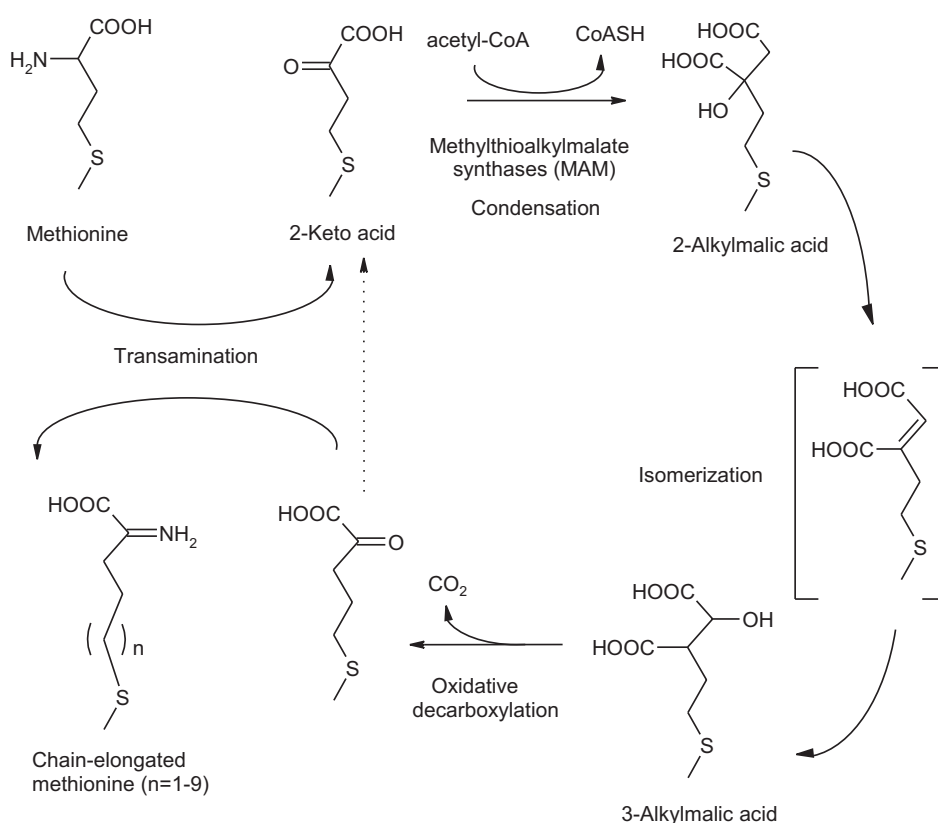
from side chain elongation of the amino acid precursors prior to the formation of the glucosinolate core structure and from a wide range of secondary modifications including oxidation, desaturation, hydroxylation, methoxylation, sulfation and glucosylation (Table 1) (1, 2).

Plants accumulating glucosinolates always possess a  $\beta$ -thioglucosidase (EC 3.2.3.1) called myrosinase, which catalyzes the hydrolysis of glucosinolates to numerous compounds with diverse biological activities. The enzyme only comes into contact with its glucosinolate substrates if the plant tissues are disrupted as a result of wounding, insect or pathogen attack. The chemical nature of hydrolysis products depends mainly on the structure of the glucosinolate side chain, plant species and reaction conditions (3, 4). Glucosinolates and their hydrolysis products are frequently studied as plant defense system against insects, herbivores and certain microbial pathogens. Besides, they serve as attractants to specialist insects feeding on crucifers (5). Mostly volatile hydrolysis products are responsible for characteristic taste and smell of cruciferous vegetables. In some *Brassica* vegetables such as cauliflower, Brussels sprouts, cabbage and broccoli, glucosinolate degradation products, especially isothiocyanates have been shown to have anticarcinogenic properties (6). However, the presence of degradation products is not always beneficial. For instance, the amount of rape meal that can be used in animal food supplement is restricted due to the goitrogenic effect of 5-vinyloxazolidine-2-thione, the spontaneous cyclization product of 2-hydroxy-3-butenyl glucosinolate which accounts for up to 80% of total glucosinolates in rape seed (7). Diverse biological properties of glucosinolates and their hydrolysis products are the reason why these plant secondary metabolites attract the interest of researchers coming from different research fields. Rapid development of molecular and genetic tools in combination with the availability of new data on the model plant *Arabidopsis thaliana* has greatly enhanced the gain of knowledge in recent years. Nevertheless, there are still many unanswered questions e.g. how glucosinolate diversity and accumulation are regulated in detail. Further research is necessary to allow precise glucosinolate manipulation in order to exploit the potential of these compounds in improving pest resistance, health and nutritional value of crop plants. This review summarizes the recent progress in glucosinolate biosynthesis, degradation and organization. Furthermore, current knowledge on the potential role of the glucosinolate-myrosinase system in plants is discussed.

**TABLE 1**

Side chain structure of some glucosinolates. *R* denotes the general structure of glucosinolate.

Glucosinolate	Trivial name	Side chain structure
2-Propenyl	Sinigrin	
3-Butenyl	Gluconapin	
3-Hydroxypropyl	–	
4-Hydroxybutyl	–	
3-Methylsulfinylpropyl	Glucoiberin	
4-Methylsulfinylbutyl	Glucoaphanin	
5-Methylsulfinylpentyl	Glucoalyssin	
6-Methylsulfinylhexyl	Glucohesperin	
7-Methylsulfinylheptyl	Glucoibarin	
8-Methylsulfinyloctyl	Glucohirsutin	
3-Methylthiopropyl	Glucoibervirin	
4-Methylthiobutyl	Glucoerucin	
6-Methylthiohexyl	Glucoquerellin	
7-Methylthioheptyl	–	
8-Methylthiooctyl	–	
Indol-3-ylmethyl	Glucoibassin	
4-Methoxyindol-3-ylmethyl	4-Methoxyglucoibassin	
1-Methoxyindol-3-ylmethyl	Neoglucobrassicin	
4-Hydroxyindol-3-ylmethyl	4-Hydroxyglucoibassin	
Benzyl	Glucoptacolin	
2-Phenylethyl	Gluconastrutiin	



**Figure 2.** The chain elongation pathway of methionine in glucosinolate biosynthesis (1).

### Glucosinolate biosynthesis

The biosynthesis of glucosinolates comprises three phases: (i) amino acid chain elongation, in which additional methylene groups are inserted into the side chain, (ii) conversion of the amino acid moiety to the glucosinolate core structure, and (iii) subsequent side chain modifications.

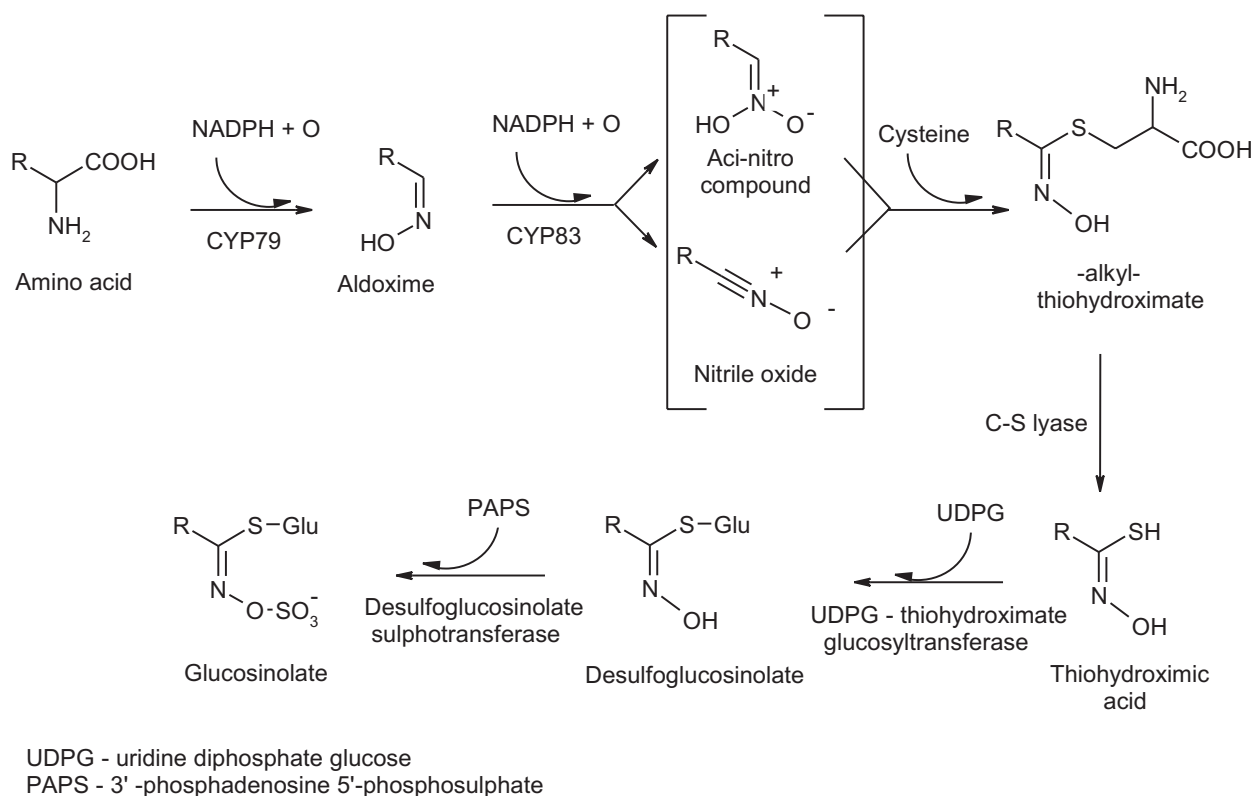
#### Side chain elongation

The majority of glucosinolates in *A. thaliana* as well as in many other species of the *Brassicaceae* are formed from methionine that has been modified by the sequential addition of 1-9 additional methylene groups to its side chain (Figure 2) (8). The pathway is initiated by transamination of methionine to form the corresponding 2-oxo acid, which is then extended by one methylene group in a three-step cycle consisting of condensation with acetyl-CoA, isomerization and oxidative decarboxylation. The newly formed 2-oxo acid can be transformed to the corresponding methionine derivative or can undergo further cycles of chain elongation. The pathway is similar to the single methylene group incorporation that occurs in leucine biosynthesis (1).

The first step, the condensation reaction, is considered to be critical for side chain variation. Quantitative trait loci (QTL) mapping and fine-scale mapping in *Arabidopsis*, using recombinant inbred lines (RIL) derived from a cross of the ecotypes Landsberg *erecta* (rich

in homomethionine-propylglucosinolates) and Columbia (rich in dihomomethionine-butylglucosinolates) identified the *Gsl-elong* locus at chromosome V (9, 10). Based on amino acid similarity to known isopropylmalate synthases (IPMS) which are responsible for condensation reaction in leucine biosynthesis, four *A. thaliana* candidate genes potentially encoding the condensing enzyme of the elongation cycle were identified. Two of these genes are at chromosome I (Atlg74040, Atlg8500). Their deduced amino acid sequences share about 90% identity with each other and are more than 60% identical to other plant IPMSs. The other two (At5g23010 and At5g23020) display lower identity to known plant IPMSs but share 85% identity with each other and show an identical intron/exon structure (10). In *A. thaliana* ecotype Columbia the latter two genes were identified as methylthioalkylmalate synthases (*MAM*) and have been shown to be responsible for the condensation step of the chain elongation cycle. At5g23010 was designated methylthioalkylmalate synthase 1 (*MAMI*) and At5g23020 as methylthioalkylmalate synthase-3 (*MAM3*). Both enzymes have similar properties but differ in their substrate specificity. *MAMI* catalyzes the condensation reaction only of the first three elongation cycles, while *MAM3* enzyme is capable to process all six additions of methylene groups (11, 12).

The remaining two steps of the chain elongation cycle (isomerization and oxidative decarboxylation) have not



**Figure 3.** Biosynthesis of glucosinolate core structure (1).

been characterized though candidate genes have been suggested based on coexpression analysis (13). The enzymatic steps are presumed to be homologous with the parallel reactions in leucine biosynthesis.

### Biosynthesis of glucosinolate core structure

The conversion of the precursor amino acid to the final glucosinolate involves intermediates common to all glucosinolates (Figure 3). The two initial steps of core glucosinolate biosynthetic pathway are catalyzed by cytochrome P450 enzymes belonging to the *CYP79* and *CYP83* families, respectively.

Cytochrome P450 enzymes of the *CYP79* family catalyze the conversion of amino acids to corresponding aldoximes. Five of the seven functional *CYP79* homologues found in the *Arabidopsis* genome have been characterized. The substrate of *CYP79A2* is phenylalanine (1), *CYP79B2* and *CYP79B3* convert tryptophan to indole-3-acetaldoxime (14, 15), and *CYP79F1* and *CYP79F2* metabolize chain-elongated methionine derivatives (16, 17, 18). *CYP79F1* and *CYP79F2* differ in their substrate specificity. *CYP79F1* is able to metabolize all chain-elongated methionine derivatives, whereas the catalytic activity of *CYP79F2* is restricted to pentahomo- and hexahomomethionine (18).

The second step in glucosinolate formation generates an unstable *aci*-nitro intermediate that conjugates with

the thiol group of cysteine via the  $\alpha$ -carbon atom. This step is catalyzed by members of the *CYP83* family. *CYP83A1* and *CYP83B1* have been identified in *Arabidopsis* (16, 19, 20). Biochemical characterization of recombinant *CYP83A1* and *CYP83B1* shows that both enzymes can metabolize all the aldoximes tested. However, *CYP83A1* has a high affinity for aliphatic aldoximes, whereas *CYP83B1* prefers indole-3-acetaldoxime and aromatic aldoximes as substrates (20, 21).

The remaining steps of glucosinolate biosynthesis involve enzymes that are thought to accommodate nearly all glucosinolate precursors regardless of their side chain. The *S*-alkylthiohydroximates are converted to thiohydroxamic acids in a reaction catalyzed by a C-S lyase, which was recently identified using a bioinformatics approach in *A. thaliana* (22). Enzymes for glycosylation (uridine diphosphate thiohydroximate glucosyltransferase) and sulfation (3'-phosphoadenosine 5'-phosphosulfate: desulfoglucosinolate sulfotransferase) have been characterized in several crucifers and partially purified (2, 23).

### Secondary modifications

Following the formation of the basic glucosinolate structure, a wide range of modifications can occur at the side chain as well as at the glucose moiety. These modifications include oxidation, hydroxylation, methoxylation, desaturation, sulfation and glycosylation, and take place in an organ and development-specific pattern (24, 25).

Based on genetic studies, a model for the side-chain modification of aliphatic glucosinolates, which are most extensively modified, has been proposed (26). Three genetic loci have been shown to be involved. The *Gsl-oxid* locus controls the oxidation of methylthio- to methylsulfanylalkylglucosinolates. With the flavin-monoxygenase GS-OX1, the appropriate S-oxygenating enzyme has been identified recently (27). The *Gsl-alk* locus controls the removal of the methylsulfanyl residue and the introduction of a double bond and, the *Gsl-oh* locus is responsible for the hydroxylation of butenylglucosinolate. In *Arabidopsis*, an additional locus termed *Gsl-ohp* controls the conversion of methylsulfanylpropyl- to hydroxypropylglucosinolate (11). Kliebenstein et al. identified three potential genes, *AOP1*, *AOP2* and *AOP3*, which all encode 2-oxoglutarate-dependent dioxygenases. No function was assigned to *AOP1*. Heterologous expression of genes in *E. coli* showed that *AOP2* catalyzes the conversion of 3-methylsulfanylpropyl- and 4-methylsulfanylbutylglucosinolate to the corresponding alkenylglucosinolates, while *AOP3* can convert 3-methylsulfanylpropyl- to 3-hydroxypropylglucosinolate (28, 29).

### Glucosinolate degradation

The plants which are able to synthesize glucosinolates always also possess a  $\gamma$ -thioglucosidase known as myrosinase (EC 3.2.3.1). The loss of cellular integrity as a result of wounding, insect or pathogen attack activates the binary glucosinolate-myrosinase system and leads to the generation of thioglucose, sulfate and an unstable intermediate which rearranges spontaneously into several degradation products (3, 4). Chemical conditions such as pH, availability of ferrous ions and presence of myrosinase-interacting proteins determine the final composition of the product mix which can include isothiocyanates, oxazolidine-2-thiones, nitriles, epithionitriles, and thiocyanates (Figure 4).

Hydrolysis at neutral conditions typically results in the formation of isothiocyanates. If a hydroxyl group at the C-2 of the glucosinolate side chain is present, the isothiocyanates formed are unstable and cyclize to oxazolidine-2-thiones. At acidic pH and in the presence of  $\text{Fe}^{2+}$  ions, the formation of nitriles occurs *in vitro* (30), while *in vivo* a protein factor such as epithiospecifier protein (ESP) is involved (31, 32). In the presence of ESP, glucosinolates with a terminal double bond in their side chain have an epithionitrile as hydrolysis product. Thiocyanates are exclusively formed from benzyl-, allyl-, and 4-methylsulfanylbutyl-glucosinolates (33). Thiocyanate-forming protein (TFP) has been shown to be involved in their formation (34). Indole glucosinolate breakdown products differ from others due to the instability of the initially formed isothiocyanates at neutral or slightly acidic pH resulting in the production of indole-methanols, ascorbic acid conjugates, and oligomeric mixtures (35).

Myrosinase activity is carried out by a group of isoenzymes present in all *Brassicaceae* species examined and is also found in 14 other plant families. Myrosinase is a dimeric protein with a molecular weight in the range of 62–75 kDa *per* subunit. Purified and characterized enzymes have shown to be highly glycosylated and are characterized by varying degrees of ascorbic acid activation. Distribution of myrosinase isoenzymes seems to be both organ-specific and species-specific (3, 4). Most myrosinases hydrolyze multiple glucosinolate substrates, but some are highly specific (32, 36). The substrate specificity can be affected by associated factors like epithiospecifier protein, myrosinase binding protein, myrosinase binding protein-related protein, and myrosinase associated proteins, although the role of these proteins has not yet been clarified (4, 31, 37, 38, 39).

Enzymes with myrosinase activity have also been found in fungi, such as *Aspergillus sydowi* (40) and *Aspergillus*

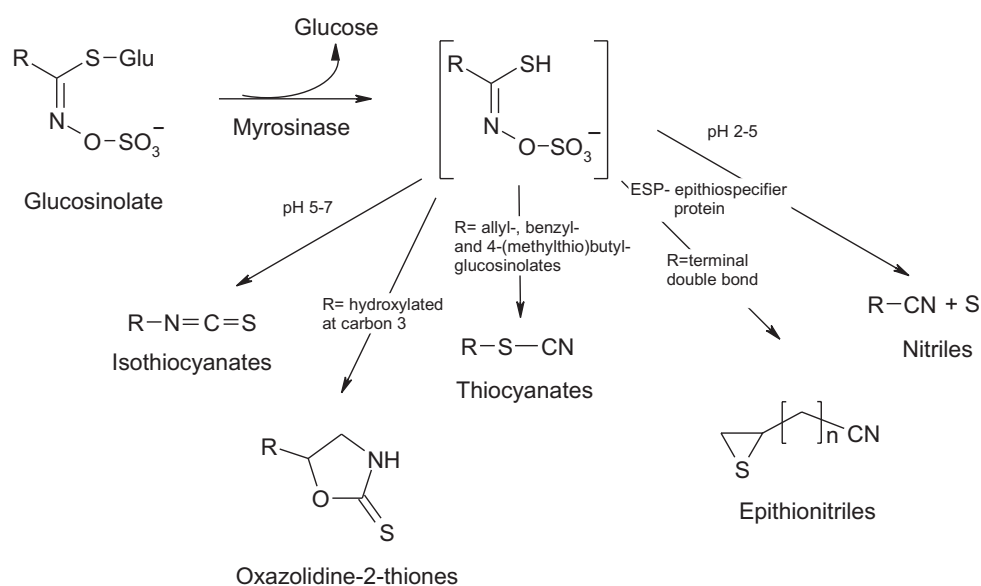


Figure 4. Structure of possible glucosinolate degradation products (1).



storage pool which can be mobilized through hydrolysis by myrosinases (65). Glucosinolates also contain nitrogen and the glucosinolate-myrosinase system can be considered to be a sink for both nutrients. It was expected that fertilizers would have an influence on this system in *Brassica* crops (4). Zhao et al. showed a clear influence of both nitrogen and sulfur supply on glucosinolates in *B. napus* (66). Their results suggest that an increase in nitrogen supply favors the hydrolysis step converting 3-butenyl to (2*R*)-2-hydroxy-3-butenyl glucosinolates. Aliphatic glucosinolates show a greater sensitivity to sulfur deficiency than indole glucosinolates due to the already sulfur containing precursor methionine. Several studies have shown that increased sulfur availability increased the glucosinolate content (61, 62), whereas a decreasing sulfur supply resulted in a decrease of free sulfate and glucosinolates while at the same time myrosinase activity increased (65, 67). Later studies demonstrated that sulfate was the main storage compound in the vegetative tissue (68) and sulfur measurements in *B. napus* showed that glucosinolates contained only a small portion of the crop's total sulfur (69). More recently, a combination of metabolite and transcript profiling revealed coordinated repression of most glucosinolate pathway genes in response to sulfate limitation (70). Taken all together, the relation between sulfur metabolism and the glucosinolate-myrosinase system exists, but it is not likely that glucosinolates are a major source of recyclable sulfur.

### Glucosinolate metabolism in growth regulation

Indole glucosinolates have been proposed as precursors for the plant hormone indole-3-acetic acid (IAA). The indole glucosinolate are supposedly hydrolyzed to indole acetonitrile (IAN), which could be hydrolyzed further to IAA by nitrilase (71, 72). Recently, several studies have provided evidence for a link between indole glucosinolates and IAA. Indole-3-acetaldoxime, the first intermediate in the indole glucosinolate biosynthesis and a product of the reaction catalyzed by CYP79B2 and CYP79B3, was found to be a precursor of IAA and is considered to be the branch point between the two metabolic pathways (20). Plants that overexpressed CYP79B2 displayed elevated levels of indole glucosinolates and IAA, while *Arabidopsis cyp79B2 cyp79B3* double mutants were strongly deficient in indole glucosinolates and partially deficient in IAA, suggesting the existence of another pathway for IAA production (73). In contrast to, this *Arabidopsis* mutants for the genes involved in the late steps of biosynthesis of indole glucosinolates all display high levels of IAA and a corresponding dwarf phenotype (including adventitious root). Those mutants further demonstrate that disruption of the conversion of indole-3-acetaldoxime to indole glucosinolates causes increased flux into IAA (20, 22). Based on structural similarity, it was also found that the indole phytoalexin camelexin is synthesized directly from indole-3-acetaldoxime (74). Thus, in *Arabidopsis* indole-3-acetaldoxime represents a key metabolic branching point at which the flow of indole-3-acetaldoxime into the biosynthetic pathways of

indole glucosinolates, camalexin and IAA must be tightly regulated (Figure 5).

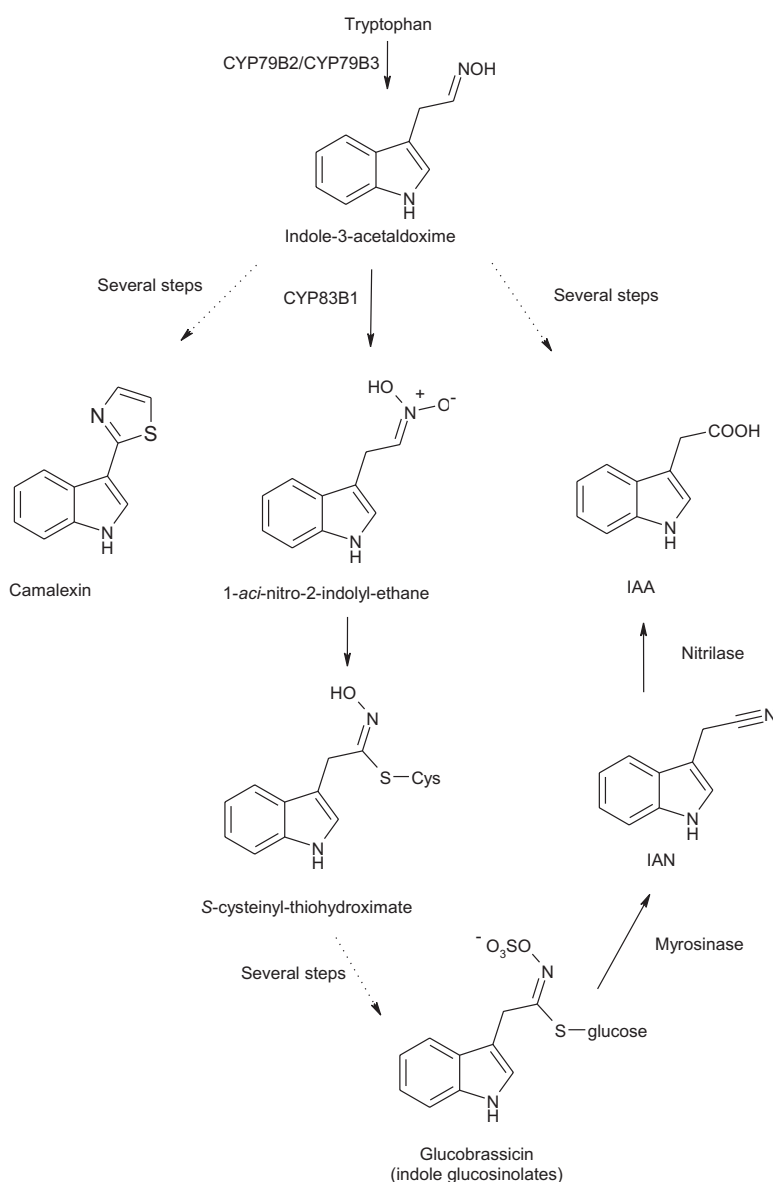
Analysis of an *Arabidopsis* CYP79F1 knockout mutant showed that abolishing the formation of short-chain methionine-derived glucosinolates is accompanied by increased levels of IAA and cytokinin (17, 63, 64). Although there is no evidence for CYP79F1 involvement in hormone metabolism, several explanations have been proposed. It is likely that the accumulation of high levels of short-chain elongated methionine and derivatives perturbs the methionine metabolism. Amongst others, methionine is the precursor for *S*-adenosylmethionine which is involved in many biosynthetic processes (i.e. ethylene biosynthesis, *trans*-methylation and regulation of cytokinin oxidase) (18).

It has also been reported that unilateral blue light promotes myrosinase activity in the illuminated side of *Raphanus sativa* hypocotyls, leading to the degradation of 4-methylthio-3-butenyl glucosinolates into natural growth inhibitors, the raphanusanins, which were involved in phototropic growth while the levels of IAA were uninfected (60). Other researchers doubted the existence of natural raphanusanins and claimed them to be artifacts of the extraction method (75). So, the possible interaction between glucosinolate metabolism, especially aliphatics, and plant development is still unclear and opens a field for further research.

### The role of the glucosinolate-myrosinase system in plant-insect/herbivore interactions

Glucosinolates and their hydrolysis products clearly play a role as mediators in plant-insect interactions. In general, plant-feeding insects can be classified as generalists or specialists. The role of the glucosinolate-myrosinase system consequently differs, i.e. glucosinolates can serve as general poison and deterrent for generalists while at the same time they can attract and stimulate feeding and egg laying of insects which are specialists on cruciferous plants.

Several studies have reported that glucosinolates exhibit growth inhibition or feeding deterrence to a wide range of general herbivores such as birds, slugs and generalist insects (26, 76). It was also found that plants respond to herbivore or insect damage by systematically accumulating higher levels of glucosinolates and thus presumably increasing their resistance (77). Usually it is the indole glucosinolates which become induced. Mewis et al. studied the glucosinolate response in *A. thaliana* to phloem-feeding aphids, the generalist *Myzus persicae* and the specialist *Brevicoryne brassicae*, and to a generalist caterpillar species *Spodoptera exigua* Hubner. They report an increase in short-chain aliphatic methylsulfinyl glucosinolates for all three insect species as well as in long-chain 8-methylsulfinyloctyl glucosinolate (8MSOO) which increased only in response to *S. exigua*. Surprisingly, indole glucosinolates were not significantly affected (78). In a field experiment using lines of *Brassica napus* which differed in their glucosinolate content,



**Figure 5.** Indole-3-acetaldoxime represents a key metabolic branching point between indole glucosinolates, camalexin and IAA biosynthesis (20, 74)

Giamoustaris and Mithen (1995) found that increasing levels of glucosinolates resulted in a decrease of damage by generalist herbivores. The opposite was true for *Brassicaceae* specialists. Furthermore, it was found that a decrease in the side chain length of aliphatic glucosinolates and in the extent of hydroxylation increased the feeding amount by *Psylliodes chrysocephala*, suggesting that this *Brassicaceae* specialist is more responsive to particular types of glucosinolates than to the others (76).

It has been recognized and accepted that glucosinolates serve as cues for feeding and oviposition of many insect herbivores which have become specialists on glucosinolate-containing plant (79, 80). Volatile hydrolysis products may serve as a signal for attraction from a distance, whereas intact glucosinolates might act as contact cues for feeding or oviposition stimulation. Behavioral

experiments were confirmed by electrophysiological investigations in which receptor organs or cells responded directly to glucosinolates or their hydrolysis products (80, 81). Volatiles produced by glucosinolates can also attract natural enemies of herbivores such as parasitoids and provide indirect protection of the plant (82, 83).

Herbivores that specialize on glucosinolate-containing plants have a mechanism to overcome the toxicity of glucosinolates and their hydrolysis products. For instance, *Plutella xylostella* possesses its own sulfatase gut enzyme that removes the sulfate moiety from the glucosinolate structure. The resulting desulfoglucosinolate does not serve as a substrate for myrosinase anymore and passes through the insect's digestive tract (84). *Pieris rapae* uses nitrile-specifying protein to direct hydrolysis toward nitriles which are less toxic than the usually produced



isothiocyanates (5). Another way of specialist herbivore protection is the possibility to sequester glucosinolates in their own tissues without harm and exploit them in self-protection. Several examples of glucosinolate-sequestering insects have been recently described, including the harlequin bug, *Murgantia histrionica* (85), the sawfly, *Athalia rosae* (86) and the aphids, *Brevicoryne brassicae* and *Lipobis erysimi* (87). To ensure the normal function of glucosinolates in defense, myrosinase should be present and therefore glucosinolate-sequestering insects contain endogenous myrosinase or rely on the myrosinase activity present in the guts of their enemies. The aphid *Brevicoryne brassicae* has been reported to possess its own myrosinase (88) which produces isothiocyanates from sequestered glucosinolates when the aphid is damaged or killed. Interestingly, these isothiocyanates even serve as alarm signal to other members of the colony (87).

### The role of the glucosinolate-myrosinase system in plant/pathogen interactions

The role of glucosinolates in defense against pathogens is less clear than that for herbivores. There are many reports demonstrating the toxicity of glucosinolate hydrolysis products to bacteria and fungi *in vitro* (89, 90), but only a few *in vivo* studies were able to correlate glucosinolates with pathogen resistance (91).

*Brassica* crops were recognized as »break crops« due to glucosinolates and their hydrolysis products which show inhibitory effects on soil borne pathogens (92, 93). Field experiments showed that wheat grown after Indian mustard and canola gives greater yield than if grown after wheat (94). Also, it was reported that the concentration of isothiocyanates achievable in the soil from the breakdown of *Brassica* tissue was sufficient to control pathogenic fungi (92). In addition the dominant soil fungal species found near glucosinolate-containing crops differ from the normal species found elsewhere and show increased tolerance to isothiocyanates (95). The hydrolysis products of indole glucosinolates were demonstrated to stimulate growth of certain ectomycorrhizal species (96).

When the relative antifungal activity of several isothiocyanate breakdown products from different glucosinolates was compared, it was found that aromatic isothiocyanates were more toxic than aliphatic ones and that the fungal toxicity of aliphatic isothiocyanates decreased with increasing length of the side chain (97, 98). Tiernes *et al.* studied the antimicrobial role of crude aqueous extracts from *Arabidopsis*. In one of the fractions, 4-methylsulphinylbutyl isothiocyanate was identified as a major compound with a broad spectrum of antimicrobial activity. A wide range of the fungi and bacteria tested showed 50% inhibition *in vitro* at a concentration lower than 350  $\mu$ M of 4-methylsulphinylbutyl isothiocyanate (99). When they tested the resistance of an *Arabidopsis* MAM1 mutant which after damage exhibits a lower amount of 4-methylsulphinylbutyl isothiocyanate, only *Fusarium oxysporum* was found to be significantly more aggressive than on wild-type plants, suggesting that glucosinolate-derived isothiocyanates might play

a role in the protection of *Arabidopsis* against particular pathogens (99).

Development of clubroot disease of the *Brassicaceae*, caused by the obligate biotrophic *Plasmodiophora brassicae*, was related to an increase of auxin and cytokinins resulting in increased cell division and cell elongation (100, 101). The high IAA content was attributed to the conversion of indole glucosinolates to IAN by myrosinase, and further conversion by nitrilase to IAA (102, 103). Ludwig-Müller *et al.* (1999) investigated several *Arabidopsis* mutants altered in glucosinolate levels. They could not draw a general conclusion about the role of indole glucosinolates in the clubroot disease development. However, almost all mutants as well as Columbia wild type showed an increase of indole-3-methylglucosinolate, and some of the mutants showed less susceptibility to the pathogen. In this study, the levels of certain aliphatic glucosinolates, 2-hydroxy-3-butenyl and 3-butenyl, were also increased after infection in both wild type Columbia and some mutants, although the authors did not discuss that point (103).

Among all studies, there were few investigating both changes of glucosinolates in inoculated leaves (= local response), and in non-inoculated leaves (= systemic response) (103, 104). Doughty *et al.* (1991) studied the glucosinolate response after *Alternaria brassicae* infection of two *Brassica napus* cultivars, Bienvenu (low in erucic acid) and Cobra (low in erucic acid and glucosinolates). Different cultivars showed different glucosinolate changes after inoculation. Local induction was more obvious in the Bienvenu cultivar, aliphatic glucosinolates accumulated rapidly but declined later while indole and aromatic glucosinolates accumulated at a slower rate and reached a maximum after 16 days. In a similar manner an induction of indole and aromatic glucosinolates were found in the Cobra cultivar. A systemic response with a threefold increase of aliphatic glucosinolates was found only in the Cobra cultivar 5 days after inoculation (104). Li *et al.* (1999) studied both local and systemic changes in glucosinolate pattern after *Sclerotinia sclerotiorum* inoculation of several *Brassica napus* cultivars. In the same way, as previously reported, different cultivars showed different glucosinolate pattern both locally and systemically. Changes were mostly due to the increase of indole and aromatic glucosinolates, although an increase of aliphatic glucosinolates was also reported (91).

In analogy to herbivores, certain pathogens may have evolved to be able to tolerate and detoxify glucosinolates. For example, high levels of 4-pentenyl glucosinolate did not correlate with resistance against *Leptosphaeria maculans* (105), perhaps due to the existence of a hydratase which is able to degrade nitriles to the less toxic formamides (106).

### Glucosinolates and defense signal molecules

During their lifetime, plants have to react to various threats coming from the environment, including micro-

bial pathogens and herbivores. As a plant is unable to escape an attack by moving to a more favorable environment, plants have evolved a broad range of defense mechanisms. The main signal molecules, recognized by the plant, are derivatives of jasmonic acid, salicylic acid and ethylene which mediate the plant response resulting in the activation of distinct sets of defense genes (107, 108). Several studies indicated changes in glucosinolate pattern after treatment with signal molecules providing another indication for a defense related role of glucosinolates. Previous studies showed that exogenous JA application usually elicits dramatic increase of indole glucosinolates while the aliphatic ones remain unchanged (109, 110). For instance, the treatment of white mustard and oilseed rape with methyl jasmonate (MeJA) accumulated indole-3-ylmethyl glucosinolate (109). In other studies, oilseed rape and *Arabidopsis* treated with MeJA were shown to accumulate both indole-3-ylmethyl glucosinolate and *N*-methoxyindole-3-ylmethyl glucosinolate (104). By contrast, SA application to the roots increased all classes of glucosinolates in the shoots of oilseed rape, with aromatic glucosinolate levels increasing more than those of indole and aliphatic ones (111).

Mikkelsen *et al.* studied glucosinolate content in response to either a single or a combination of treatments with MeJA, 2,6-dichloro-isonicotinic acid (INA) as functional homologue of SA and 1-aminocyclopropane-1-carboxylate (ACC) as ethylene precursor in *A. thaliana* (112). The results obtained confirmed the accumulation of indole glucosinolates after MeJA treatment, e.g., the concentration of all indole glucosinolates increased upon treatment with MeJA. Treatment with INA or ACC induced 4-methoxyindole-3-ylmethyl glucosinolate levels. Treatments with both MeJA and ACC induced indole-3-ylmethyl and *N*-methoxyindole-3-ylmethyl glucosinolate levels while accumulation of 4-methoxyindole-3-ylmethyl glucosinolate was absent. Changes of aliphatic glucosinolate levels were also noticed. Upon MeJA treatment the levels of 5-methylsulfinylpentyl, 8-methylthiooctyl and 8-methylsulfinyloctyl glucosinolates increased significantly while the levels of 8-methylsulfinyloctyl glucosinolates also increased after INA, ACC and both MeJA and ACC treatments (112).

Kliebenstein *et al.* report altered glucosinolate accumulation upon MeJA and SA treatment of different ecotypes, suggesting different responses in different *A. thaliana* cultivars. After treatment of the Landsberg ecotype Ler with MeJA and SA, four different glucosinolate response patterns were identified which were not seen in the ecotype Columbia Col-0. The first pattern is characterized by a MeJA induction and includes the 3-hydroxypropyl, indole-3-ylmethyl and 1-methoxyindole-3-ylmethyl glucosinolates. The MeJA induction maximized within 24 hours, after which the levels of 3-hydroxypropyl glucosinolate returned to control levels, while indole-3-ylmethyl glucosinolate levels remained elevated. The second response pattern was the induction of 4-methoxyindole-3-ylmethyl glucosinolate as the only glucosinolate induced exclusively by SA but not by MeJA. The third re-

sponse pattern was a synergistic action of both MeJA and SA which resulted in the accumulation of 8-methylsulfinyloctyl glucosinolate after 48 hours. The last response included 8-methylthiooctyl glucosinolate that showed no regulation by any of the compounds tested (113).

A recent interesting study has monitored the response of glucosinolates in *Brassica oleracea* and *Brassica nigra* after the application of JA and SA either on shoots or on roots. There were changes in all classes of glucosinolates, but those changes differed depending on the treatment (root or shoot application of signal molecules) as well as on the species (114). For example, *B. oleracea* after JA treatment of either shoot or root or both showed increased levels of all indole and some aliphatic glucosinolates in shoots. In roots, a similar increase was observed in case of aliphatic glucosinolates while indoles were not affected. The treatment of root and both root and shoot with SA induced some aliphatic glucosinolates only in shoot, while the treatment with both SA and JA induced both aliphatic and indole glucosinolates also in shoots (114).

From all the glucosinolate literature present, it is evident that the overall defense response of a plant is a rather complex process which considerably varies depending on environmental conditions and the species involved and therefore we are far from a complete understanding of glucosinolate function in plant defense. Numerous studies have shown that glucosinolates and their hydrolysis products have an impact on herbivores and pathogens; however, many questions about the mechanism of glucosinolate induction, the signaling pathways involved and the plant's potential benefit from glucosinolate accumulation remain to be solved.

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## REFERENCES

1. WITTSTOCK U, HALKIER B A 2002 Glucosinolate research in the *Arabidopsis* era. *Trends Plant Sci* 7: 263–270
2. HALKIER B A, GERSHENZON J 2006 Biology and biochemistry of glucosinolates. *Annu Rev Plant Biol* 57: 303–333
3. BONES A M, ROSSITER J T 1996 The myrosinase-glucosinolate system, its organisation and biochemistry. *Physiol Plant* 97: 194–208
4. RASK L, ANDREASSON E, EKBOM B, ERIKSSON S, PONTOPPIDAN B, MEIJER J 2000 Myrosinase: gene family evolution and herbivore defense in *Brassicaceae*. *Plant Mol Biol* 42: 93–113
5. WITTSTOCK U, AGERBIRK N, STAUBER E J, OLSEN C E, HIPPLER M, MITCHELL-OLDS T, GERSHENSON J, VOGEL H 2004 Successful herbivore ontack due to metabolic diversion of a plant chemical defense. *Proc Natl Acad Sci U S A* 101: 4859–4864
6. FAHEY J W, ZALCMANN A T, TALALAY P 2001 The chemical diversity and distribution of glucosinolates and isothiocyanates among plants. *Phytochemistry* 56: 5–51
7. CHEN S, ANDREASSON E 2001 Update on glucosinolate metabolism and transport. *Plant Physiol Biochem* 39: 743–758

8. GRASER G, SCHNEIDER B, OLDHAM N J, GERSHENZON J 2000 The methionine chain elongation pathway in the biosynthesis of glucosinolates in *Eruca sativa* Brassicaceae. *Arch Biochem Biophys* 378: 411–419
9. MITHEN R, CAMPOS H 1996 Genetic variation of aliphatic glucosinolates in *Arabidopsis thaliana* and prospects for map based gene cloning. *Entomol Exp Appl* 80: 202–205
10. KROYMANN J, TEXTORS, TOKUHISA J G, FALK K L, BARTRAM S, GERSHENZON J, MITCHELL-OLDS T 2001 A gene controlling variation in *Arabidopsis* glucosinolate composition is part of the methionine chain elongation pathway. *Plant Physiol* 127: 1077–1088
11. TEXTOR S, BARTRAM S, KROYMANN J, FALK K L, HICK A, PICKETT J A, GERSHENZON J 2004 Biosynthesis of methionine-derived glucosinolates in *Arabidopsis thaliana*: recombinant expression and characterization of methylthioalkylmalate synthase, the condensing enzyme of the chain-elongation cycle. *Planta* 218: 1026–1035
12. TEXTOR S, DE KRAKER W, HAUSE B, GERSHENZON J, TOKUHISA J G 2007 MAM3 Catalyzes the Formation of All Aliphatic Glucosinolate Chain Lengths in *Arabidopsis*. *Plant Physiol* 144: 60–71
13. HIRAI MY, SUGIYAMA K, SAWADA Y, TOHGE T, OBAYASHI T, SUZUKI A, ARAKI R, SAKURAI N, SUZUKI H, AOKI K, GODA H, NISHIZAWA O I, SHIBATA D, SAITO K 2007 Omics-based identification of *Arabidopsis* Myb transcription factors regulating aliphatic glucosinolate biosynthesis. *Proc Natl Acad Sci U S A* 104: 6478–6483
14. HULL A K, VIJ R, CELENZA J L 2000 *Arabidopsis* cytochrome P450s that catalyze the first step of tryptophan-dependent indole-3-acetic acid biosynthesis. *Proc Natl Acad Sci U S A* 97: 2379–2384
15. MIKKELSEN M D, HANSEN C H, WITTSTOCK U, HALKIER B A 2000 Cytochrome P450 CYP79B2 from *Arabidopsis* catalyzes the conversion of tryptophan to indole-3-acetaldoxime, a precursor of indole glucosinolates and indole-3-acetic acid *J Biol Chem* 275: 33712–33717
16. HANSEN C H, DU L C, NAUR P, OLSEN C E, AXELSEN K B, HICK A J, PICKETT J A, HALKIER B A 2001 CYP83B1 is the oxime-metabolizing enzyme in the glucosinolate pathway in *Arabidopsis*. *J Biol Chem* 276: 24790–24796
17. REINTANZ B, LEHNEN M, REICHELT M, GERSHENZON J, KOWALCZYK M, SANDBERG G, GODDE M, UHL R, PALME K 2001 *Bus*, a bushy *Arabidopsis* CYP79F1 knockout mutant with abolished synthesis of short-chain aliphatic glucosinolates. *Plant Cell* 13: 351–367
18. CHEN S X, GLAWISCHNIG E, JORGENSEN K, NAUR P, JORGENSEN B, OLSEN C E, HANSEN C H, RASMUSSEN H, PICKETT J A, HALKIER B A 2003 CYP79F1 and CYP79F2 have distinct functions in the biosynthesis of aliphatic glucosinolates in *Arabidopsis*. *Plant J* 33: 923–937
19. BAK S, OLSEN C E, PETERSEN B L, MOLLER B L, HALKIER B A 1999 Metabolic engineering of p-hydroxybenzylglucosinolate in *Arabidopsis* by expression of the cyanogenic CYP79A1 from *Sorghum bicolor*. *Plant J* 20: 663–671
20. BAK S, TAX F E, FELDMANN K A, GALBRAITH D W, FEYERREISEN R 2001 CYP83B1, a cytochrome P450 on the metabolic branch point in auxin and indole glucosinolate biosynthesis in *Arabidopsis*. *Plant Cell* 13: 101–111
21. NAUR P, PETERSEN B L, MIKKELSEN M D, BAK S, RASMUSSEN H, OLSEN C E, HALKIER B A 2003 CYP83A1 and CYP83B1, two nonredundant cytochrome P450 enzymes metabolizing oximes in the biosynthesis of glucosinolates in *Arabidopsis*. *Plant Physiol* 133: 63–72
22. MIKKELSEN M D, NAUR P, HALKIER B A 2004 *Arabidopsis* mutants in the C-S lyase of glucosinolate biosynthesis establish a critical role for indole-3-acetaldoxime in auxin homeostasis. *Plant J* 37: 770–777
23. GRUBB C D, ZIPP B J, LUDWIG-MULLER J, MASUNO M N, MOLINSKI T F, ABEL S 2004 *Arabidopsis* glucosyltransferase UGT74B1 functions in glucosinolate biosynthesis and auxin homeostasis. *Plant J* 40: 893–908
24. PETERSEN B L, CHEN S X, HANSEN C H, OLSEN C E, HALKIER B A 2002 Composition and content of glucosinolates in developing *Arabidopsis thaliana*. *Planta* 214: 562–571
25. BROWN P D, TOKUHISA J G, REICHELT M, GERSHENZON J 2003 Variation of glucosinolate accumulation among different organs and developmental stages of *Arabidopsis thaliana*. *Phytochemistry* 62: 471–481
26. GIAMOUSTARIS A, MITHEN R 1996 Genetics of aliphatic glucosinolates 4 Side-chain modification in *Brassica oleracea*. *Theor Appl Genet* 93: 1006–1010
27. HANSEN B G, KLIEBENSTEIN D J, HALKIER B A 2007 Identification of a flavin-monoxygenase as the S-oxygenating enzyme in aliphatic glucosinolate biosynthesis in *Arabidopsis*. *Plant J* 50: 902–910
28. KLIEBENSTEIN D J, GERSHENZON J, MITCHELL-OLDS T 2001 Comparative quantitative trait loci mapping of aliphatic, indolic and benzylic glucosinolate production in *Arabidopsis thaliana* leaves and seeds. *Genetics* 159: 359–370
29. KLIEBENSTEIN D J, KROYMANN J, BROWN P, FIGUTH A, PEDERSEN D, GERSHENZON J, MITCHELL-OLDS T 2001 Genetic control of natural variation in *Arabidopsis* glucosinolate accumulation. *Plant Physiol* 126: 811–825
30. UDA Y, KURATA T, ARAKAWA N 1986 Effects of pH and ferrous ion on the degradation of glucosinolates by myrosinase. *Agr Biol Chem* 50: 2735–2740
31. LAMBRIX V, REICHELT M, MITCHELL-OLDS T, KLIEBENSTEIN D J, GERSHENZON J 2001 The *Arabidopsis* epithiospecifier protein promotes the hydrolysis of glucosinolates to nitriles and influences *Trichoplusia ni* herbivory. *Plant Cell* 13: 2793–2807
32. BERNARDI R, FINIGUERRA M G, ROSSI A A, PALMIERI S 2003 Isolation and biochemical characterization of a basic myrosinase from ripe *Crambe abyssinica* seeds, highly specific for epi-progoitrin. *J Agric Food Chem* 51: 2737–2744
33. HASAPIS X, MACLEOD A J 1982 Benzylglucosinolate degradation in heat-treated *Lepidium sativum* seeds and detection of a thiocyanate-forming factor. *Phytochemistry* 21: 1009–1013
34. BUROW M, BERGNER A, GERSHENZON J, WITTSTOCK U 2007 Glucosinolate hydrolysis in *Lepidium sativum*-identification of the thiocyanate-forming protein. *Plant Mol Biol* 63: 49–61
35. AGERBIRK N, OLSEN C E, SORENSEN H 1998 Initial and final products, nitriles, and ascorbigenes produced in myrosinase-catalyzed hydrolysis of indole glucosinolates. *J Agric Food Chem* 46: 1563–1571
36. CHEN S X, HALKIER B A 1999 Functional expression and characterization of the myrosinase MYR1 from *Brassica napus* in *Saccharomyces cerevisiae*. *Protein Expr Purif* 17: 414–420
37. TAIPALENSUU J, FALK A, RASK L 1996 A wound- and methyl jasmonate-inducible transcript coding for a myrosinase-associated protein with similarities to an early nodulin. *Plant Physiol* 110: 483–491
38. ZABALA M D, GRANT M, BONES A M, BENNETT R, LIM Y S, KISSEN R, ROSSITER J T 2005 Characterisation of recombinant epithiospecifier protein and its over-expression in *Arabidopsis thaliana*. *Phytochemistry* 66: 859–867
39. BUROW M, MARKERT J, GERSHENZON J, WITTSTOCK U 2006 Comparative biochemical characterization of nitrile-forming proteins from plants and insects that alter myrosinase-catalysed hydrolysis of glucosinolates. *FEBS J* 273: 2432–2446
40. REESE E T, CLAPP R C, MANDELS M 1958 A thioglucosidase in fungi. *Arch Biochem Biophys* 75: 228–242
41. OHTSURU M, TSURUO I, HATA T 1973 The production and stability of intracellular myrosinase from *Aspergillus niger*. *Agr Biol Chem* 37: 967–971
42. TANI N, OHTSURU M, HATA T 1974 Isolation of myrosinase producing microorganism. *Agr Biol Chem* 38: 1617–1622
43. OGINSKY E L, STEIN A E, GREER M A 1965 Myrosinase activity in bacteria as demonstrated by the conversion of progoitrin to goitrin. *Proc Soc Exp Biol Med* 119: 360–364
44. GOODMAN I, FOUTS J R, BRESNICK E, MENEGAS R, HITCHINGS G H 1959 A mammalian thioglycosidase. *Science* 130: 450–451
45. JONES A M, BRIDGES M, BONES A M, COLE R, ROSSITER J T 2001 Purification and characterisation of a non-plant myrosinase from the cabbage aphid *Brevicoryne brassicae* (L.). *Insect Biochem Mol Biol* 31: 1–5.

46. KAZANA E, POPE T W, TIBBLES L, BRIDGES M, PICKETT J A, BONES A M, POWELL G, ROSSITER J T 2007 The cabbage aphid: a walking mustard oil bomb. *Proc Biol Sci* 274: 2271–2277.
47. DU L C, HALKIER B A 1998 Biosynthesis of glucosinolates in the developing silique walls and seeds of *Sinapis alba*. *Phytochemistry* 48: 1145–1150
48. CHEN S X, PETERSEN B L, OLSEN C E, SCHULZ A, HALKIER B A 2001 Long-distance phloem transport of glucosinolates in *Arabidopsis*. *Plant Physiol* 127: 194–201
49. KOROLEVA O A, DAVIES A, DEEKEN R, THORPE M R, TOMOS A D, HEDRICH R 2000 Identification of a new glucosinolate-rich cell type in *Arabidopsis*. flower stalk *Plant Physiol* 124: 599–608
50. FALK K L, VOGEL C, TEXTOR S, BARTRAM S, HICK A, PICKETT J A, GERSHENZON J 2004 Glucosinolate biosynthesis: demonstration and characterization of the condensing enzyme of the chain elongation cycle in *Eruca sativa*. *Phytochemistry* 65: 1073–1084
51. LÜTHY B, MATILE P 1984 The mustard oil bomb – rectified analysis of the subcellular organization of the myrosinase system. *Biochemie Und Physiologie Der Pflanzen* 179: 5–12
52. THANGSTAD O P, IVERSEN T H, SLUPPHAUG G, BONES A 1990 Immunocytochemical localization of myrosinase in *Brassica napus* L. *Planta* 180: 245–248
53. THANGSTAD O P, EVJEN K, BONES A 1991 Immunogold-EM localization of myrosinase in *Brassicaceae*. *Protoplasma* 161: 85–93
54. HOGLUND A S, LENMAN M, RASK L 1992 Myrosinase is localized to the interior of myrosin grains and is not associated to the surrounding tonoplast membrane. *Plant Sci* 85: 165–170
55. ANDREASSON E, JORGENSEN L B, HOGLUND A S, RASK L, MEIJER J 2001 Different myrosinase and idioblast distribution in *Arabidopsis* and *Brassica napus*. *Plant Physiol* 127: 1750–1763
56. HUSEBYE H, CHADCHAWAN S, WINGE P, THANGSTAD O P, BONES A M 2002 Guard cell- and phloem idioblast-specific expression of thioglucoside glucohydrolase 1 myrosinase in *Arabidopsis*. *Plant Physiol* 128: 1180–1188
57. THANGSTAD O P, GILDE B, CHADCHAWAN S, SEEM M, HUSEBYE H, BRADLEY D, BONES A M 2004 Cell specific, cross-species expression of myrosinases in *Brassica napus*, *Arabidopsis thaliana* and *Nicotiana tabacum*. *Plant Mol Biol* 54: 597–611
58. GROB K, MATILE P 1979 Vacuolar location of glucosinolates in horseradish root-cells. *Plant Sci L* 14: 327–335
59. MATILE P 1980 The mustard oil bomb – compartmentation of the myrosinase system. *Biochemie Und Physiologie Der Pflanzen* 175: 722–731
60. HASEGAWA T, YAMADA K, KOSEMURA S, YAMAMURA S, HASEGAWA K 2000 Phototropic stimulation induces the conversion of glucosinolate to phototropism-regulating substances of radish hypocotyls. *Phytochemistry* 54: 275–279
61. KAUR S, GUPTA S K, SUKHIJA P S, MUNSHI S K 1990 Accumulation of glucosinolates in developing mustard *Brassica juncea* L seeds in response to sulfur application. *Plant Sci* 66: 181–184
62. UNDERHILL E W, CHISHOLM M D, STECK W 1980 E-5, Z-7-Dodecadienal, a sex-pheromone component of the western tent caterpillar, *Malacosoma californicum* Lepidoptera, Lasiocampidae *Can Entomol* 112: 629–641
63. TANTIKANJANA T, MIKKELSEN M D, HUSSAIN M, HALKIER B A, SUNDARESAN V 2004 Functional analysis of the tandem-duplicated P450 genes SPS/BUS/CYP79F1 and CYP79F2 in glucosinolate biosynthesis and plant development by Ds transposon-generated double mutants. *Plant Physiol* 135: 840–848
64. TANTIKANJANA T, YONG J W H, LETHAM D S, GRIFFITH M, HUSSAIN M, LJUNG K, SANDBERG G, SUNDARESAN V 2001 Control of axillary bud initiation and shoot architecture in *Arabidopsis* through the SUPERSHOOT gene. *Genes Dev* 15: 1577–1588
65. SCHNUG E, CEYNOWA J 1990 Phytopathological aspects of glucosinolates in oilseed rape. *J Agron Crop Sci-Z Acker Pflanzenbau* 165: 319–328
66. MAILER R J 1989 Effects of applied sulfur on glucosinolate and oil concentrations in the seeds of rape *Brassica napus* L and turnip rape *Brassica rapa* L var silvestris lam briggs. *Aust J Agric Res* 40: 617–624
67. FALK K L, TOKUHISA J, GERSHENZON J 2007 The effect of sulphur nutrition on plant glucosinolate content: physiology and molecular mechanisms. *Plant Biology* 9: 573–581
68. ZHAO F J, WOOD A P, MCGRATH S P 1999 Effects of sulphur nutrition on growth and nitrogen fixation of pea *Pisum sativum* L. *Plant Soil* 212: 209–219
69. FIELDSSEND J, MILFORD G F J 1994 Changes in glucosinolates during crop development in single-low and double-low genotypes of winter oilseed rape *Brassica napus* L Production and distribution in vegetative tissues and developing pods during development and potential role in the recycling of sulfur within the crop. *Ann Appl Biol* 124: 531–542
70. HIRAI M Y, KLEIN M, FUJIKAWA Y, YANO M, GOODENOWE D B, YAMAZAKI Y, KANAYA S, NAKAMURA Y, KITAYAMA M, SUZUKI H, SAKURAI N, SHIBATA D, TOKUHISA J, REICHEL T M, GERSHENZON J, PAPANBROCK J, SAITO K 2005 Elucidation of gene-to-gene and metabolite-to-gene networks in *Arabidopsis* by integration of metabolomics and transcriptomics. *J Biol Chem* 280: 25590–25595
71. SEARLE L M, CHAMBERLAIN K, RAUSCH T, BUTCHER D N 1982 The conversion of 3-Indolylmethylglucosinolate to 3-Indolylacetonitrile by myrosinase, and its relevance to the clubroot disease of the *Cruciferae*. *J Exp Bot* 33: 935–942
72. BARTEL B, FINK G R 1994 Differential regulation of an auxin-producing nitrilase gene family in *Arabidopsis thaliana* *Proc Natl Acad Sci U S A* 91: 6649–6653
73. ZHAO Y D, HULL A K, GUPTA N R, GOSS K A, ALONSO J, ECKER J R, NORMANLY J, CHORY J, CELENZA J L 2002 Trp-dependent auxin biosynthesis in *Arabidopsis*: involvement of cytochrome P450s CYP79B2 and CYP79B3. *Genes Dev* 16: 3100–3112
74. GLAWISCHNIG E, HANSEN B G, OLSEN C E, HALKIER B A 2004 Camalexin is synthesized from indole-3-acetaldoxime, a key branching point between primary and secondary metabolism in *Arabidopsis*. *Proc Natl Acad Sci U S A* 101: 8245–8250
75. MATSUOKA H, TODA Y, YONEYAMA K, UDA Y 1998 Formation of raphanusanins depends on extraction procedure and solvent. *Phytochemistry* 47: 975–977
76. GIAMOUSTARIS A, MITHEN R 1995 The effect of modifying the glucosinolate content of leaves of oilseed rape *Brassica napus* ssp *oleifera* on its interaction with specialist and generalist pests. *Ann Appl Biol* 126: 347–363
77. MARTIN N, MÜLLER C 2006 Induction of plant responses by a sequestering insect: relationship of glucosinolate concentration and myrosinase activity. *Basic Appl Ecol* 8: 13–25
78. MEWIS I, APPEL H M, HOM A, RAINA R, SCHULTZ J C 2005 Major signaling pathways modulate *Arabidopsis* glucosinolate accumulation and response to both phloem-feeding and chewing insects. *Plant Physiol* 138: 1149–1162
79. MEWIS I Z, ULRICH C, SCHNITZLER W H 2002 The role of glucosinolates and their hydrolysis products in oviposition and host-plant finding by cabbage webworm, *Hellula undalis*. *Entomol Exp Appl* 105: 129–139
80. MILES C I, DEL CAMPO M L, RENWICK J A 2005 Behavioral and chemosensory responses to a host recognition cue by larvae of *Pieris rapae*. *J Comp Physiol Neuroethol Sens Neural Behav Physiol* 191: 147–155
81. ROJAS J C 1999 Electrophysiological and behavioral responses of the cabbage moth to plant volatiles. *J Chem Ecol* 25: 1867–1883
82. READ D P, FEENY P P, ROOT R B 1970 Habitat selection by the aphid parasite *Diaeretiella rapae* Hymenoptera: Braconidae and hyperparasite *Charips brassicae* Hymenoptera: Cynipidae. *Can Entomol* 102: 1567–1578
83. MATTIACCI L, DICKE M, POSTHUMUS M A 1994 Induction of parasitoid attracting synomone in *Brussels sprouts* plants by feeding of *Pieris brassicae* larvae – role of mechanical damage and herbivore elicitor. *J Chem Ecol* 20: 2229–2247
84. RATZKA A, VOGEL H, KLIEBENSTEIN D J, MITCHELL-OLDS T, KROYMANN J 2002 Disarming the mustard oil bomb. *Proc Natl Acad Sci U S A* 99: 11223–11228
85. ALIABADI A, RENWICK J A, WHITMAN D W 2002 Sequestration of glucosinolates by harlequin bug *Murgantia histrionica*. *J Chem Ecol* 28: 1749–1762

86. MULLER C, AGERBIRK N, OLSEN C E, BOEVE J L, SCHAFNER U, BRAKEFIELD P M 2001 Sequestration of host plant glucosinolates in the defensive hemolymp of the sawfly *Athalia rosae*. *J Chem Ecol* 27: 2505–2516
87. BRIDGES M, JONES A M E, BONES A M, HODGSON C, COLE R, BARTLET E, WALLSGROVE R, KARAPAPA V K, WATTS N, ROSSITER J T 2002 Spatial organization of the glucosinolate-myrosinase system in brassica specialist aphids is similar to that of the host plant. *Proc R Soc Lond Ser B-Biol Sci* 269: 187–191
88. JONES A M E, WINGE P, BONES A M, COLE R, ROSSITER J T 2002 Characterization and evolution of a myrosinase from the cabbage aphid *Brevicoryne brassicae*. *Insect Biochem Mol Biol* 32: 275–284
89. SMOLINSKA U, MORRA M J, KNUDSEN G R, JAMES R L 2003 Isothiocyanates produced by *Brassicaceae* species as inhibitors of *Fusarium oxysporum*. *Plant Dis* 87: 407–412
90. MARI M, LEONI O, IORI R, CEMBALI T 2002 Antifungal vapour-phase activity of allyl-isothiocyanate against *Penicillium expansum* on pears. *Plant Pathol* 51: 231–236
91. LI Y, KIDDLE G, BENNETT R N, WALLSGROVE R M 1999 Local and systemic changes in glucosinolates in Chinese and European cultivars of oilseed rape *Brassica napus* L after inoculation with *Sclerotinia sclerotiorum* stem rot. *Ann Appl Biol* 134: 45–58
92. ANGUS J F, GARDNER P A, KIRKEGAARD J A, DESMARCHELIER J M 1994 Biofumigation – isothiocyanates released from *Brassica* roots inhibit growth of the take-all fungus. *Plant Soil* 162: 107–112
93. BROWN P D, MORRA M J 1996 Hydrolysis products of glucosinolates in *Brassica napus* tissues as inhibitors of seed germination. *Plant Soil* 181: 307–316
94. KIRKEGAARD J A, GARDNER P A, ANGUS J F, KOETZ E 1994 Effect of *Brassica* break crops on the growth and yield of wheat. *Aust J Agric Res* 45: 529–545
95. ISHIMOTO H, FUKUSHI Y, YOSHIDA T, TAHARA S 2000 *Rhizopus* and *Fusarium* are selected as dominant fungal genera in rhizospheres of *Brassicaceae*. *J Chem Ecol* 26: 2387–2399
96. ZENG R S, MALLIK A U, SETLIFF E 2003 Growth stimulation of ectomycorrhizal fungi by root exudates of *Brassicaceae* plants: Role of degraded compounds of indole glucosinolates. *J Chem Ecol* 29: 1337–1355
97. MANICI L M, LAZZERI L, PALMIERI S 1997 In vitro fungitoxic activity of some glucosinolates and their enzyme-derived products toward plant pathogenic fungi. *J Agric Food Chem* 45: 2768–2773
98. SARWAR M, KIRKEGAARD J A 1998 Biofumigation potential of brassicas – II Effect of environment and ontogeny on glucosinolate production and implications for screening. *Plant Soil* 201: 91–101
99. TIERENS K, THOMMA B P H, BROUWER M, SCHMIDT J, KISTNER K, PORZEL A, MAUCH-MANI B, CAMMUE B P A, BROEKAERT W F 2001 Study of the role of antimicrobial glucosinolate-derived isothiocyanates in resistance of *Arabidopsis* to microbial pathogens. *Plant Physiol* 125: 1688–1699
100. LUDWIG-MÜLLER J, BENDEL U, THERMANN P, RUPPEL M, EPSTEIN E, HILGENBERG W 1993 Concentrations of indole-3-acetic-acid in plants of tolerant and susceptible varieties of chinese-cabbage infected with *Plasmidiophora brassicae* woron. *New Phytol* 125: 763–769
101. LUDWIG-MÜLLER J, PIEPER K, RUPPEL M, COHEN J D, EPSTEIN E, KIDDLE G, BENNETT R 1999 Indole glucosinolate and auxin biosynthesis in *Arabidopsis thaliana* L Heynh glucosinolate mutants and the development of clubroot disease. *Planta* 208: 409–419
102. BUTCHER D N, EL-TIGANI S, INGRAM D S 1974 The role of indole glucosinolates in the club root disease of the *Cruciferae*. *Physiol Plant Pathol* 4: 127–140
103. LUDWIG-MÜLLER J, SCHUBERT B, PIEPER K, IHMIG S, HILGENBERG W 1997 Glucosinolate content in susceptible and resistant Chinese cabbage varieties during development of clubroot disease. *Phytochemistry* 44: 407–414
104. DOUGHTY K J, PORTER A J R, MORTON A M, KIDDLE G, BOCK C H, WALLSGROVE R 1991 Variation in the glucosinolate content of oilseed rape *Brassica napus* L leaves 2 response to infection by *Alternaria brassicae* Berk Sacc. *Ann Appl Biol* 118: 469–477
105. ROUXEL T, KOLLMANN A, BOULIDARD L, MITHEN R 1991 Abiotic elicitation of indole phytoalexins and resistance to *Leptosphaeria maculans* within *Brassicaceae*. *Planta* 184: 271–278
106. SEXTON A C, HOWLETT B J 2000 Characterisation of a cyanide hydratase gene in the phytopathogenic fungus *Leptosphaeria maculans*. *Mol Gen Genet* 263: 463–470
107. REYMOND P, FARMER E E 1998 Jasmonate and salicylate as global signals for defense gene expression. *Curr Opin Plant Biol* 1: 404–411
108. TURNER J G, ELLIS C, DEVOTO A 2002 The jasmonate signal pathway. *Plant Cell* 14: S153–S164
109. BODNARYK R P 1994 Potent effect of jasmonates on indole glucosinolates in oilseed rape and mustard. *Phytochemistry* 35: 301–305
110. DOUGHTY K J, KIDDLE G A, PYE B J, WALLSGROVE R M, PICKETT J A 1995 Selective induction of glucosinolates in oilseed rape leaves by methyl jasmonate. *Phytochemistry* 38: 347–350
111. KIDDLE G A, DOUGHTY K J, WALLSGROVE R M 1994 Salicylic acid-induced accumulation of glucosinolates in oilseed rape *Brassica napus* L leaves. *J Exp Bot* 45: 1343–1346
112. MIKKELSEN M D, HALKIER B A 2003 Metabolic engineering of valine- and isoleucine-derived glucosinolates in *Arabidopsis* expressing CYP79D2 from cassava. *Plant Physiol* 131: 773–779
113. KLIEBENSTEIN D J, FIGUTH A, MITCHELL-OLDS T 2002 Genetic architecture of plastic methyl jasmonate responses in *Arabidopsis thaliana*. *Genetics* 161: 1685–1696
114. AN DAM N M, WITJES L, SVATOS A 2004 Interactions between aboveground and belowground induction of glucosinolates in two wild *Brassica* species. *New Phytol* 161: 801–810