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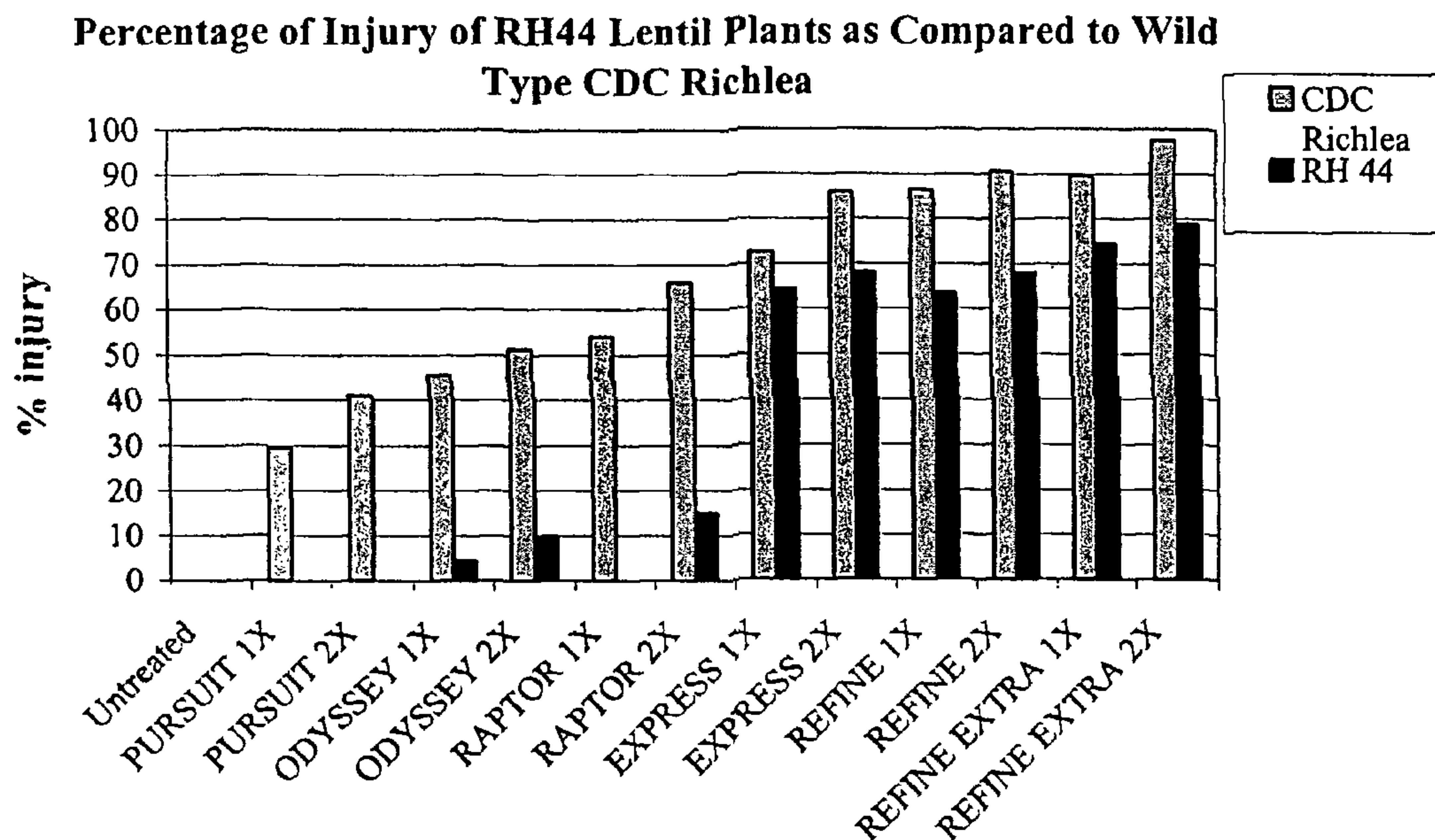
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(54) **Titre : PLANTES DE LENTILLES PRESENTANT UNE RESISTANCE ACCRUE AUX HERBICIDES IMIDAZOLINONE**
(54) **Title: LENTIL PLANTS HAVING INCREASED RESISTANCE TO IMIDAZOLINONE HERBICIDES**



(57) **Abrégé/Abstract:**

The present invention is directed to lentil plants having increased resistance to an imidazolinone herbicide. One such plant described herein is the RH44 lentil variety. The present invention also includes seeds produced by these lentil plants and methods of controlling weeds in the vicinity of these lentil plants.

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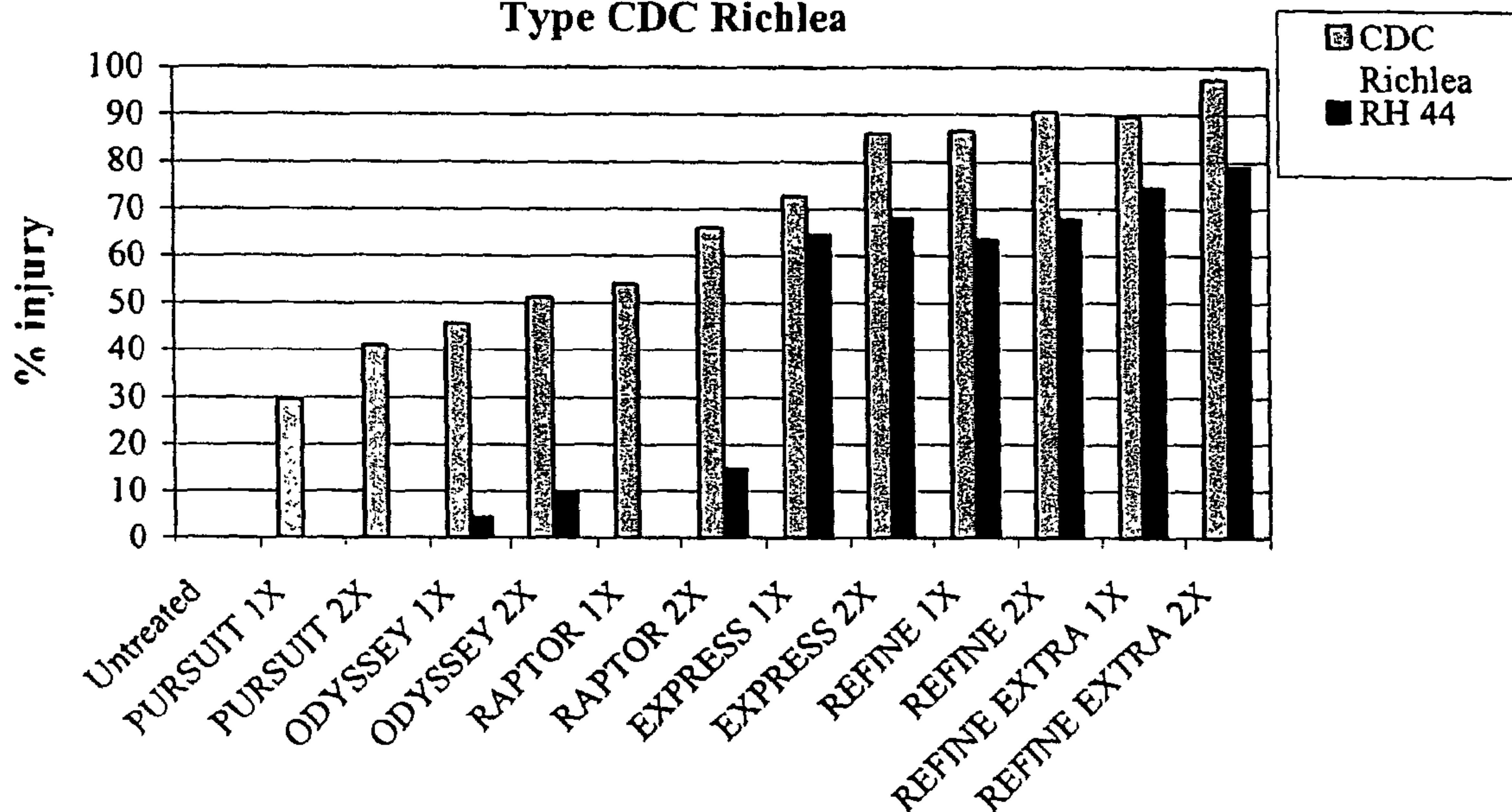
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(54) Title: LENTIL PLANTS HAVING INCREASED RESISTANCE TO IMIDAZOLINONE HERBICIDES

Percentage of Injury of RH44 Lentil Plants as Compared to Wild
Type CDC Richlea



(57) Abstract: The present invention is directed to lentil plants having increased resistance to an imidazolinone herbicide. One such plant described herein is the RH44 lentil variety. The present invention also includes seeds produced by these lentil plants and methods of controlling weeds in the vicinity of these lentil plants.

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LENTIL PLANTS HAVING INCREASED RESISTANCE TO IMIDAZOLINONE HERBICIDES

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FIELD OF THE INVENTION

The present invention relates in general to plants having an increased resistance to imidazolinone herbicides. More specifically, the present invention relates to lentil plants obtained by mutagenesis and cross-breeding that have an increased resistance to imidazolinone herbicides.

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BACKGROUND OF THE INVENTION

Imidazolinone and sulfonylurea herbicides are widely used in modern agriculture due to their effectiveness at very low application rates and relative non-toxicity in animals. Imidazolinone and sulfonylurea herbicides inhibit the activity of acetohydroxyacid synthase (AHAS), or acetolactate synthase (ALS) (E.C.4.1.3.18), the key enzyme in the biosynthesis of branch chain amino acids such as valine, leucine and isoleucine (Shaner et al. 1984 Plant Physiol. 76:545-546). By inhibiting AHAS activity, this class of herbicides prevents further growth and development of susceptible plants including many weed species. Several examples of commercially available imidazolinone herbicides are PURSUIT® (imazethapyr), SCEPTER® (imazaquin) and ARSENAL® (imazapyr). Examples of sulfonylurea herbicides are chlorsulfuron, metsulfuron methyl, sulfometuron methyl, chlorimuron ethyl, thifensulfuron methyl, tribenuron methyl, bensulfuron methyl, nicosulfuron, ethametsulfuron methyl, rimsulfuron, triflurosulfuron methyl, triasulfuron, primisulfuron methyl, cinosulfuron, amidosulfuron, fluzasulfuron, imazosulfuron, pyrazosulfuron ethyl and halosulfuron.

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Due to their high effectiveness and low-toxicity, imidazolinone herbicides are favored for application by spraying over the top of a wide area of vegetation. The ability to spray an herbicide over the top of a wide range of vegetation decreases the costs associated with plantation establishment and maintenance and decreases the need for site preparation prior to use of such chemicals. Spraying over the top of a desired tolerant species also results in the ability to achieve maximum yield potential of the desired species due to the absence of competitive species. However, the ability to use such spray-over techniques is dependent upon the presence of imidazolinone resistant species of the desired vegetation in the spray over area.

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Among the major agricultural crops, some leguminous species such as soybean are naturally resistant to imidazolinone herbicides due to their ability to rapidly metabolize the herbicide compounds (Shaner and Robinson 1985 Weed Sci. 33:469-471). Other crops such as corn (Newhouse et al. 1992 Plant Physiol. 100:882-886) and

rice (Barrette et al. 1989 Crop Safeners for Herbicides, Academic Press New York, pp. 195-220) are somewhat susceptible to imidazolinone herbicides. The differential sensitivity to the imidazolinone herbicides is dependent on the chemical nature of the particular herbicide and differential metabolism of the compound from a toxic to a non-toxic form in each plant (Shaner et al. 1984 Plant Physiol. 76:545-546; Brown et al. 1987 Petic. Biochem. Physiol. 27:24-29). Other plant physiological differences such as absorption and translocation also play an important role in sensitivity (Shaner and Robinson 1985 Weed Sci. 33:469-471).

Computer-based modeling of the three dimensional conformation of the AHAS-inhibitor complex predicts several amino acids in the proposed inhibitor binding pocket as sites where induced mutations would likely confer selective resistance to imidazolinones (Ott et al. 1996 J. Mol. Biol. 263:359-368). Lentil plants produced with these rationally designed mutations in the proposed binding sites of the AHAS enzyme have in fact exhibited specific resistance to a single class of herbicides (Ott et al. 1996 J. Mol. Biol. 263:359-368). Other mutations in the AHAS gene have been linked to resistance to the imidazolinone herbicides in canola (Swanson et al. 1989 Theor. Appl. Genet. 78:525-530) and corn (Newhouse et al. 1991 Theor. Appl. Genet. 83:65-70).

Studies of the ALS gene in other crop plants have also resulted in sulfonylurea and imidazolinone resistance in those plants. In one report, use of a mutant ALS gene from *Arabidopsis* coupled with selection on sulfonylurea herbicide resulted in the production of resistant transgenic rice plants (Li et al. 1992 Plant Cell Rep. 12:250-255). An increase in *in vitro* resistance to chlorsulfuron of similar magnitude (200-fold) was demonstrated in transgenic rice containing a 35S/ALS transgene (Li et al. 1992 Plant Cell Rep. 12:250-255), and imidazolinone-resistant growth of transgenic tobacco was 100-fold greater than non-transformed control plants (Sathasivan et al. 1991 Plant Physiol. 97:1044-1050).

Expression of the introduced AHAS or ALS gene at different magnitudes has also been achieved by manipulating several aspects of the transformation including the use of different promoters and screening larger populations of transformants (Odell et al. 1990 Plant Physiol. 94:1647-1654; Sathasivan et al. 1991 Plant Physiol. 97:1044-1050; Li et al. 1992 Plant Cell Rep. 12:250-255). Studies showed that replacing the *Arabidopsis* ALS promoter with the CaMV35S promoter resulted in 40-fold differences in *in vitro* resistance to chlorsulfuron (Li et al. 1992 Plant Cell Rep. 12:250-255). In tobacco, the increase in resistance to imazethapyr in individual calli transformed with a mutant ALS gene from *Arabidopsis* ranged from 10- to 1000-fold, most likely reflecting the differences in gene copy numbers or in chromosomal positions of the transgenes (Sathasivan et al. 1991 Plant Physiol. 97:1044-1050).

Plant resistance to imidazolinone has also been reported in a number of patents. U.S. Patent No. 4,761,373 generally describes the use of an altered AHAS gene to elicit

herbicide resistance in plants, and specifically discloses certain imidazolinone resistant corn lines. U.S. Patent No. 5,013,659 discloses plants exhibiting herbicide resistance possessing mutations in at least one amino acid in one or more conserved regions. The mutations described therein encode either cross-resistance for imidazolinones and sulfonylureas or sulfonylurea-specific resistance, but imidazolinone-specific resistance is not described. Additionally, U.S. Patent No. 5,731,180 and U.S. Patent No. 5,767,361 discuss an isolated gene having a single amino acid substitution in a wild-type monocot AHAS amino acid sequence that results in imidazolinone-specific resistance.

However, to date, the prior art has not described an imidazolinone resistant pulse crop such as lentil. Pulses are the seeds of legumes that are used as food, including pea, bean, lentil, chickpea and fababean. Pulse crops, provide about 10% of the total dietary protein of the world. Lentil was one of the earliest cultivated crops in the world with archeological evidence from the early Stone Age. Lentil remains an important source of dietary protein in India, Southwest Asia and the Mediterranean, and Canadian lentil production is primarily directed toward export to these regions. While lentil is grown mainly for the seed to be harvested as a food export, the straw can also be used as a high quality animal feed or as a source of organic material for soil improvement. Cultivated varieties of lentil (*Lens culinaris*) are believed to descend from *Lens orientalis*, the only wild-type species able to naturally cross with *Lens culinaris* and produce fully fertile progeny.

A major challenge in lentil production is weed control. Lentil seedlings are short and slow-growing in relation to many weed species and therefore compete very poorly. Effective chemical weed control is necessary for commercial viability. The ability to spray over an herbicide that kills a broad spectrum of broadleaf weeds, either as a pre-emergent spray or as a post-emergent spray, would be beneficial to lentil production. Even more advantageous would be an herbicide that also controls a broad spectrum of grassy weeds and volunteer cereals that could be applied over a broad area of lentil crops.

Therefore, what are needed in the art are lentil plants having increased resistance to herbicides such as imidazolinone and methods for controlling weed growth in the vicinity of lentil plants. Such compositions and methods would allow for the use of spray over techniques when applying herbicides to areas containing lentil plants.

SUMMARY OF THE INVENTION

The present invention relates to lentil plants having increased resistance to an imidazolinone herbicide as compared to a wild type variety of the plant. The lentil plant described herein can be any member of the *Lens* genus, including, but not limited to, *Lens culinaris* Medikus, *Lens orientalis* (Boiss.) Hand.-Maz., *Lens nigricans* (M.

Bieb.) Grand., *Lens ervoides* (Bring.) Grand., *Lens odemensis* Ladiz., *Lens lamottei* Czefranova, *Lens tomentosus* Ladiz and hybrids thereof. Additionally, the imidazolinone herbicide referred to herein can be selected from, but is not limited to, imazethapyr, imazapic, imazamox, imazaquin, imazethabenz and imazapyr.

5 In one embodiment of the present invention, the lentil plant variety is designated RH44 and has a Patent Deposit Designation Number PTA-3270. The present invention also includes a mutant, recombinant, or genetically engineered derivative of the plant with Patent Deposit Designation Number PTA-3270, any progeny of the plant with Patent Deposit Designation Number PTA-3270 and a plant
10 that is the progeny of any of these plants. In further preferred embodiments, the lentil plant also has the herbicide resistance characteristics of the plant with Patent Deposit Designation Number PTA-3270.

Included in the present invention are hybrids of the RH44 line described herein and another lentil variety including, but not limited to, CDC Richlea, CDC Robin,
15 CDC Sovereign, CDC Glamis, CDC Milestone, CDC Vantage, Eston, Laird, Spanish Brown and French Green.

In addition to lentil plants having increased resistance to imidazolinone herbicides, the present invention also encompasses plant parts, plant cells and plant seeds derived from these plants. In one embodiment, the lentil plant seed is true
20 breeding for an increased resistance to an imidazolinone herbicide as compared to a wild type variety of the lentil plant seed.

The methods of the present invention include methods for controlling weeds within the vicinity of a lentil plant, comprising applying an imidazolinone herbicide to the weeds and to the lentil plant, wherein the lentil plant has increased resistance to the
25 imidazolinone herbicide as compared to a wild type variety of the lentil plant.

The invention also seeks to provide a lentil plant cell from a lentil plant having increased resistance to an imidazolinone herbicide as compared to a wild type variety of the plant, wherein the lentil plant has an ATCC Patent Deposit Designation Number PTA-3270, and is a mutant, recombinant, or genetically engineered derivative of the
30 plant with ATCC Patent Deposit Designation Number PTA-3270. The lentil plant comprises altered acetohydroxyacid synthase (AHAS) activity when compared to a wild-type lentil plant, or is a plant that is a progeny of any of these plants, wherein the lentil plant comprises altered AHAS activity when compared to a wild-type lentil plant.

Further, the invention seeks to provide a method for controlling weeds within the vicinity of a lentil plant, comprising applying an imidazolinone herbicide to the weeds and to the lentil plant, wherein the lentil plant has increased resistance to the imidazolinone herbicide as compared to a wild type variety of the lentil plant, and wherein the lentil plant, has an ATCC Patent Deposit Designation Number PTA-3270, and is a mutant, recombinant, or genetically engineered derivative of the plant with ATCC Patent Deposit Designation Number PTA-3270 or of any progeny of the plant with ATCC Patent Deposit Designation Number PTA-3270. The lentil plant comprises altered AHAS activity when compared to a wild-type lentil plant, or is a plant that is a progeny of any of these plants, wherein the lentil plant comprises altered AHAS activity when compared to a wild-type lentil plant.

Still further the invention provides a lentil plant cell from a seed of lentil line RH44, representative seed of said line having been deposited under ATCC Patent Deposit Designation Number PTA-3270.

Further still, the invention comprehends a method for controlling weeds within the vicinity of a lentil plant, the method comprising applying an imidazolinone herbicide to the weeds and to the lentil plant, wherein the lentil plant is produced by growing a seed of lentil line RH44, representative seed of said line having been deposited under ATCC Patent Deposit Designation Number PTA-3270.

Yet further, the invention provides a method for producing a hybrid lentil seed wherein the method comprises crossing a lentil plant produced by growing a seed of lentil line RH44 with a different lentil plant and harvesting the resulting hybrid lentil seed, wherein representative seed of lentil line RH44 has been deposited under ATCC Patent Deposit Designation Number PTA-3270.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a table showing the effects of amino acid substitutions in the AHAS isoenzyme on the herbicide resistance of various plants.

Figure 2 is a graph showing the percentage injury to RH44 lentil plants and the wild type CDC Richlea lentil variety when sprayed in the seedling stage with various rates of imidazolinone or sulfonylurea herbicides.

Figure 3 is a graph showing the inhibition of AHAS activity in RH44 and CDC Richlea lentil plants by a negative feedback inhibition assay.

Figure 4 is a graph showing the inhibition of AHAS activity in RH44 and CDC Richlea lentil plants by increasing concentrations of imazapic.

Figure 5 is a graph showing the inhibition of AHAS activity in RH44 and CDC Richlea lentil plants by increasing concentrations of imazethapyr.

Figure 6 is a table showing the agronomic characteristics of RH44 lentil plants and various wild type commercial lentil varieties.

Figure 7 is a table showing the increased resistance of RH44 lentil plants to ODYSSEY herbicide as compared to various wild type commercial lentil varieties.

Figure 8 is a table showing the amino acid composition of RH44 lentil plants and various wild type commercial lentil varieties.

5 Figure 9 is a table showing the nutritional composition of RH44 lentil plants as compared to other registered commercial lentil varieties.

DETAILED DESCRIPTION

10 The present invention is directed to lentil plants, lentil plant parts and lentil plant cells having increased resistance to imidazolinone herbicides. In one embodiment, a wild type lentil plant is one which is a member of the *Lens* genus and does carry the dominant gene for resistance to an imidazolinone herbicide. The present invention also includes seeds produced by the lentil plants described herein and methods for controlling weeds in the vicinity of the lentil plants described herein. It is
15 to be understood that as used in the specification and in the claims, "a" or "an" can mean one or more, depending upon the context in which it is used. Thus, for example, reference to "a cell" can mean that at least one cell can be utilized.

As used herein, the term "lentil plant" refers to a plant that is a member of the *Lens* genus of the *Leguminosae* family. The lentil plants of the present invention can be
20 members of the *Lens* genus including, but not limited to, *Lens culinaris* Medikus, *Lens orientalis* (Boiss.) Hand.-Maz., *Lens nigricans* (M. Bieb.) Grand., *Lens ervoides* (Bring.) Grand., *Lens odemensis* Ladiz., *Lens lamoittie* Czefranova and *Lens tomentosus* Ladiz. (Ladizinsky et al. 1984, van Oss et al., 1997) and hybrids thereof. The term "lentil plant" is intended to encompass lentil plants at any stage of maturity or
25 development as well as any tissues or organs taken or derived from any such plant unless otherwise clearly indicated by context. Plant tissues and organs include, but are not limited to, leaves, seeds, stems, flowers, roots, single cells, gametes, anther cultures, calli cultures, tissue cultures and protoplasts. In particular, the present invention includes seeds produced by the lentil plants of the present invention. In one
30 embodiment, the seeds are true breeding for an increased resistance to an imidazolinone herbicide as compared to a wild type variety of the lentil plant seed.

The present invention describes a lentil plant having increased resistance to an imidazolinone herbicide as compared to a wild type variety of the plant. The Examples below provide a detailed description of the mutagenesis, breeding and selection of
35 lentil plants having such increased resistance to an imidazolinone herbicide. One plant derived from these procedures is deposited with the ATCC (Patent Deposit Designation Number PTA-3270) and designated herein as the RH44 lentil variety. A deposit of 2500 seeds of the RH44 lentil variety was made with the American Type Culture Collection, Manassas, Virginia on March 20, 2001. This deposit was made in
40 accordance with the terms and provisions of the Budapest Treaty relating to the deposit

of microorganisms. The deposit was made for a term of at least thirty years and at least five years after the most recent request for the furnishing of a sample of the deposit is received by the ATCC. The deposited seeds were accorded Patent Deposit Designation Number PTA-3270.

5 The present invention includes the lentil plant having a Patent Deposit Designation Number PTA-3270; a mutant, recombinant, or genetically engineered derivative of the plant with Patent Deposit Designation Number PTA-3270; any progeny of the plant with Patent Deposit Designation Number PTA-3270; and a plant that is the progeny of any of these plants. In a preferred embodiment, the lentil plant of
10 the present invention additionally has the herbicide resistance characteristics of the plant with Patent Deposit Designation Number PTA-3270.

 Also included in the present invention are hybrids of the RH44 line described herein and another lentil variety including, but not limited to, CDC Richlea, CDC Robin, CDC Sovereign, CDC Glamis, CDC Milestone, CDC Vantage, Eston, Laird,
15 Spanish Brown and French Green. The term “variety” refers to a group of plants within a species that share constant characters that separate them from the typical form and from other possible varieties within that species. While possessing at least one distinctive trait, a variety is also characterized by some variation between individuals within the variety, based primarily on the Mendelian segregation of traits among the
20 progeny of succeeding generations. A variety is considered “true breeding” for a particular trait if it is genetically homozygous for that trait to the extent that, when the true-breeding variety is self-pollinated, a significant amount of independent segregation of the trait among the progeny is not observed. In the present invention, the trait arises from a dominant mutation in an AHAS gene of the lentil plant or seed.

25 In one embodiment of the present invention, the lentil plant having increased resistance to an imidazolinone herbicide comprises an altered AHAS nucleic acid. As used herein, the term “altered AHAS nucleic acid” refers to an AHAS nucleic acid that is mutated from an AHAS nucleic acid in a wild type lentil plant and that confers increased imidazolinone resistance to a plant in which it is transcribed. In a preferred
30 embodiment, the altered AHAS nucleic acid comprises a serine to asparagine amino acid substitution. In a more preferred embodiment, the altered AHAS nucleic acid comprises a serine to asparagine amino acid substitution in an AHAS gene. In a still further preferred embodiment, the serine to asparagine amino acid substitution corresponds to the serine to asparagine amino acid substitutions found in other AHAS
35 gene paralogs that display imidazolinone resistance. Examples of such mutated AHAS gene paralogs include those found in wheat variety SWP965001, corn variety XI-12 and canola variety PM1, all of which have increased resistance to imidazolinone herbicides and are described in Figure 1. By “AHAS nucleic acid” is meant a RNA or
40 DNA sequence that encodes or directs the expression of an AHAS protein, and may include a coding region and its corresponding untranslated 5’ and 3’ sequence regions.

Additionally, "AHAS gene" refers specifically to a DNA sequence that encodes or directs the expression of an AHAS protein.

It is to be understood that the lentil plant of the present invention can comprise a wild type or unaltered AHAS gene in addition to an altered AHAS gene. As described in Example 3, it is contemplated that the mutation in lentil variety RH44 contains a mutation in only one of two AHAS isoenzymes. Therefore, the present invention includes a lentil plant comprising one or more altered AHAS nucleic acids.

As also used herein, the term "AHAS protein" refers to an acetohydroxyacid synthase protein and the term "altered AHAS protein" refers to any AHAS protein that is mutated from a wild type AHAS protein and that confers increased imidazolinone resistance to a plant, plant cell, plant part, plant seed or plant tissue when it is expressed therein. The imidazolinone herbicide can be selected from, but is not limited to, PURSUIT® (imazethapyr), CADRE® (imazapic), RAPTOR® (imazamox), SCEPTER® (imazaquin), ASSERT® (imazethabenz), ARSENAL® (imazapyr) ODYSSEY® (imazapyr/imazamox), or a derivative thereof.

In addition to the compositions of the present invention, the present invention provides a method of controlling weeds growing in the vicinity of the lentil plants described above. These methods comprise applying imidazolinone herbicides to weeds in the vicinity of lentil plants having an increased resistance to an imidazolinone herbicide as compared to a wild type variety of the plant. In a preferred embodiment, the lentil plant comprises a altered AHAS nucleic acid. In a more preferred embodiment, the altered AHAS nucleic acid comprises a serine to asparagine amino acid substitution in an AHAS gene. In a still further preferred embodiment, the serine to asparagine amino acid substitution corresponds to the serine to asparagine amino acid substitutions found in other AHAS gene paralogs that display imidazolinone resistance.

As described above, the present invention teaches compositions and methods for increasing the imidazolinone resistance of a lentil plant or seed as compared to a wild-type variety of the plant or seed. In a preferred embodiment, the imidazolinone resistance of a lentil plant or seed is increased such that the plant or seed can withstand an imidazolinone herbicide application of preferably approximately 1-28 ounces, more preferably approximately 3-14 ounces, and most preferably approximately 6, 7, or 8 ounces of active ingredient per acre.

By providing for lentil plants having increased resistance to imidazolinone, a wide variety of formulations can be employed for protecting lentil plants from weeds, so as to enhance plant growth and reduce competition for nutrients. An imidazolinone herbicide can be used by itself for post-emergence control of weeds in areas surrounding the lentil plants described herein or an imidazolinone herbicide formulation can be used that contains other additives. Such additives include other herbicides, detergents, adjuvants, spreading agents, sticking agents, stabilizing agents,

or the like. The imidazolinone herbicide formulation can be a wet or dry preparation and can include, but is not limited to, flowable powders, emulsifiable concentrates and liquid concentrates. The imidazolinone herbicide and herbicide formulations can be applied in accordance with conventional methods, for example, by spraying, irrigation, dusting, or the like.

It should be understood that the foregoing relates to preferred embodiments of the present invention and that numerous changes may be made therein without departing from the scope of the invention. The invention is further illustrated by the following examples, which are not to be construed in any way as imposing limitations upon the scope thereof. On the contrary, it is to be clearly understood that resort may be had to various other embodiments, modifications, and equivalents thereof, which, after reading the description herein, may suggest themselves to those skilled in the art without departing from the spirit of the present invention and/or the scope of the appended claims. Additionally, all references cited herein may be referred to for further details.

EXAMPLES

EXAMPLE 1

Mutagenesis of Mixed Lentil Seed and Selection of RH44 Lentil Variety Having Increased Resistance to Imidazolinone Herbicides

Lentil line RH44 was derived from a bulk of mixed F₃ lentil (*Lens culinaris*) seed developed via conventional crossing followed by self-pollination. Five kilograms of bulk, mixed seed were treated in EMS solution and then planted in the field as the M₁ generation. M₂ seeds were harvested from the field-grown M₁ plants. M₂ plants were advanced to the M₃ generation. M₃ seed was planted on 1 hectare (ha). Prior to flowering, the field was sprayed with 2X ODYSSEY® herbicide. At harvest, approximately 300 surviving plants were harvested and threshed. Approximately 150 plants were selected for further evaluation by planting a sample of seed from each plant in pots in growth rooms. These plants were sprayed with 2X ODYSSEY® herbicide at four weeks after emergence. Eight pots were selected for field evaluation. Seed of each line was sown in a small field plot and then sprayed with 2X ODYSSEY®. Line RH44 was selected as having imidazolinone resistance derived from a population of approximately one million M₃ seeds. Since the original mutagen dose was small, the M₃ population size was large, and the modification rate was relatively low (1:150,000), it is unlikely that multiple modifications occurred in RH44 to contribute to any deleterious effects (Konzak, 1987 Induced mutations in wheat improvement. In: Heyne, E.G. (ed.) Wheat and Wheat Improvement. American Society of Agronomy, Madison, WI. pp. 428-443). Figure 2 shows the increased imidazolinone resistance of the RH44 line as compared to the CDC Richlea lentil variety. Figure 2 also shows that

the RH44 line has little resistance to sulfonyleureas such as EXPRESS, REFINE and REFINE EXTRA. The results in Figure 2 reflect tolerance readings taken two weeks after application of the herbicide. The designations "1X" and "2X" refer to commercial application rates of those products.

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EXAMPLE 2

Analysis of Mutation in Lentil Variety RH44

The available data strongly indicates that a mutation in a single gene similar to that observed in wheat, corn and canola is responsible for the observed imidazolinone tolerance in the RH44 variety of lentil. In wheat, the AHAS isozymes have been labeled as genes A, B and C. The mutation responsible for imidazolinone tolerance in wheat is due to a point mutation of AGC to AAC at a single site in AHAS gene A (Figure 1). This site corresponds to the maize amino acid position 621 previously identified in X1-12 (imidazolinone-tolerant maize). This single nucleotide change of a guanine to an adenine results in a single amino acid change serine to asparagine (Figure 1). Additionally, the imidazolinone tolerance trait in CLEARFIELD® canola is controlled by two semi-dominant genes PM1 and PM2. Resistance to the PM1 mutant is conferred through a guanine to adenine point mutation in the AHAS1 gene. This results in a similar single amino acid change from serine to asparagine. A different point mutation is responsible for the resistance observed in the PM2 mutant. A guanine to threonine change in AHAS3 results in a tryptophan to leucine amino acid change.

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The serine to asparagine amino acid substitution results in a plant that is tolerant to imidazolinone herbicides, but not sulfonyleureas (Newhouse et al. 1992 Plant Physiol. 100:882-886). This is true of wheat, corn and canola. Similarly, the RH44 lentil variety is similarly not tolerant to sulfonyleurea herbicides, suggesting that a similar guanine to adenine point mutation produced a serine to asparagine amino acid substitution similar to the other CLEARFIELD® varieties X1-12, corn, PM1 canola and SWP965001 wheat (Figure 1).

EXAMPLE 3

Herbicide Resistance of the RH44 Lentil Variety

The responses of the RH44 lentil variety and the CDC Richlea variety were identical in a negative feedback inhibition assay. As described earlier, AHAS is an enzyme active in the biosynthesis of leucine and valine. Under conditions of excess leucine and/or valine, the activity of AHAS is diminished. The ability of leucine and valine addition to inhibit AHAS activity was the same in the RH44 lentil and CDC Richlea lentil varieties, indicating very similar levels of expression and a lack of any discernible difference in AHAS function (Figure 3).

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The AHAS activity in the wild type CDC Richlea lentil variety was inhibited by imazapic (CADRE®) (Figure 4) and imazethapyr (PURSUIT®) (Figure 5) representative of all imidazolinone herbicides. The AHAS activity in the RH44 lentil variety was also inhibited by the addition of the two imidazolinone herbicides, but not to the same extent as for the wild type lentil CDC Richlea.

These results suggest the existence of a semi-dominant AHAS isozyme in the RH44 lentil variety that is encoded by genes similar to those in X1-12 corn, SWP965001 wheat and PM1 canola. It is therefore likely that the gene primarily responsible for AHAS activity in the RH44 lentil has undergone a point mutation similar to that observed in corn, wheat and canola. This has made the isozyme coded for by this sequence resistant to imidazolinone inhibition. The wild type isozyme has no such mutation and therefore is susceptible to imidazolinone herbicide. Such a scenario would explain why AHAS activity drops 30 to 40% with the addition of maximum concentrations of imidazolinone herbicides. The resistance trait was determined to be stable in the mutagenized line RH44 lentil variety indicating that the resistance trait is semi-dominant, also similar to the imidazolinone resistant trait in canola.

EXAMPLE 4

Agronomic Characteristics of the RH44 Lentil Variety

The agronomic traits (yield, height, time to maturity) of the RH44 lentil variety were similar to most of the commercial wild type cultivars of lentil. The time to flowering was approximately 58 days, the time to maturity was approximately 100 days and the mean height was 30 to 35 cm (Figure 6). Both the RH44 lentil variety and CDC Richlea lentil variety are relatively high yielding, however, they are both susceptible to ascochyta blight and anthracnose.

Initial field evaluation of RH44 lentil indicates that the seed yield is similar to that of CDC Richlea and superior to some commercial varieties (Figure 7). In field trials, the RH44 lentil has commercially viable phenotypic and agronomic properties. Harvested RH44 lentil seeds were mixed in appearance. Some seeds had a 'mottled' seed coat while some remained clear. Selection of seeds on the basis of appearance resulted in a mixed harvest in subsequent generations, which is a phenotypic trait similar to the Eston lentil variety.

EXAMPLE 5

Amino Acid Composition of Imidazolinone-tolerant Lentil

An analysis of the amino acid composition was conducted to compare the RH44 lentil variety with existing registered wild type lentil varieties in order to determine if any significant differences existed (Figure 8). The seed for analysis was obtained from trials conducted in the year 2000 where RH44 lentil and several other lentil varieties were grown under the same conditions. Amino acid composition was determined using

AOAC Method 982.30 D,E,F by Woodson-Tenent Laboratories, Inc. The results of this analysis demonstrated the similarity in amino acid composition among all varieties of lentil, many of which have very different phenotypic characteristics. This analysis also demonstrated that the mutation leading to imidazolinone tolerance in lentil produces no change outside the range of natural variability in the amino acid composition of lentils.

EXAMPLE 6

Nutritional Analysis of Imidazolinone-tolerant Lentil

A proximate analysis was conducted to compare the RH44 lentil variety with existing registered wild type lentil varieties in order to determine if any significant differences in nutritional value existed (Figure 9). The seed for analysis was obtained from trials conducted in the year 2000 where RH44 lentil and several other lentil varieties were grown under the same conditions. The results of this analysis demonstrate the similarity in many nutritional characteristics among varieties of lentil, several of which have very different phenotypic characteristics. This analysis also demonstrates that the mutation leading to imidazolinone tolerance in RH44 lentil produces no subsequent change in the moisture, fat, fiber or protein content of lentil seed. The RH44 lentil is changed in its AHAS activity, but is unchanged in nutritional/food safety attributes when compared to commercial wild type lentil varieties in Canada.

WHAT IS CLAIMED IS:

1. A cell of a lentil plant, wherein the lentil plant is of lentil line RH44, representative seed of said line having been deposited under ATCC Patent Deposit Designation Number PTA-3270, said lentil plant cell having increased resistance to an imidazolinone herbicide as compared to a cell of wild type variety of the plant.
2. A cell of a hybrid lentil plant, wherein said hybrid lentil plant is produced by the process of crossing a lentil plant of lentil line RH44, a representative seed of said line having been deposited under ATCC Patent Deposit Designation Number PTA-3270, said lentil plant having increased resistance to an imidazolinone herbicide as compared to a cell of a wild type variety of the plant, with another lentil plant to produce a hybrid lentil plant, said cell of the hybrid lentil plant comprising the increased resistance to an imidazolinone herbicide exhibited by said lentil plant of line RH44.
3. A cell of a progeny lentil plant, wherein said progeny lentil plant is produced by the process of crossing of a lentil plant of lentil line RH44, a representative seed of said line having been deposited under ATCC Patent Deposit Designation Number PTA-3270, said lentil plant having increased resistance to an imidazolinone herbicide as compared to a cell of a wild type variety of the plant, with another lentil plant to produce a hybrid lentil plant, said hybrid lentil plant used in a process to produce progeny lentil plants, said cell of a progeny lentil plant comprising the increased resistance to an imidazolinone herbicide exhibited by said lentil plant of line RH44.
4. Use of a lentil plant comprising the lentil plant cell of claim 1 for producing progeny lentil plant, wherein the progeny lentil plant comprises the increased resistance to an imidazolinone herbicide exhibited by the lentil line RH44.
5. Use of a lentil plant comprising the lentil plant cell of any one of claims 1, 2, or 3 for producing seed.
6. Use of seed from a lentil plant comprising the lentil plant cell of any one of claims 1, 2, or 3 for growing a lentil plant.

7. Use of a lentil plant comprising the lentil plant cell of any one of claims 1, 2, or 3 to cross with another lentil plant.
8. Use of a lentil plant comprising the lentil plant cell of any one of claims 1, 2, or 3 for crossing with another *Lens culinaris* variety, to produce progeny retaining the increased imidazolinone resistance characteristics of lentil line RH44.
9. Use of a lentil plant cell of any one of claims 1, 2, or 3 as a recipient of a transgene.
10. Use of a lentil plant comprising the lentil plant cell of any one of claims 1, 2, or 3 for growing a crop or producing seed.
11. Use of a progeny lentil plant comprising the lentil plant cell of any one of claims 1, 2, or 3 for growing a crop or producing seed.
12. A method for producing lentil plants, comprising:
 - a. mutating cells of any one of claims 1, 2, or 3 to produce mutant cells or preparing recombinant or genetically engineered cells from cells as claimed in any one of claims 1, 2, or 3; and
 - b. regenerating lentil plants from said mutant, recombinant or genetically engineered cells.
13. A method of growing a lentil crop or producing lentil seed comprising:
 - a. growing a progeny lentil plant comprising the lentil plant cell of any one of claims 1, 2, or 3, at a locus; and
 - b. treating said locus with an imidazolinone herbicide.
14. A method for controlling weeds in a field, comprising:
 - a. growing a progeny lentil plant comprising the lentil plant cell of any one of claims 1, 2, or 3, at a locus in said field, and
 - b. treating said locus with an imidazolinone herbicide;
 thereby controlling the weeds.

15. Use of a progeny lentil plant or a seed thereof; said plant comprising the lentil plant cell of any one of claims 1, 2, or 3; in combination with the imidazolinone herbicide for controlling weeds in an agricultural field comprising: applying an imidazolinone herbicide to the site in the field in which said progeny lentil plant is grown or said seed is planted.
16. The method of claim 13 or 14, wherein said imidazolinone herbicide comprises at least one of: imazethapyr, imazapic, imazamox, imazaquin, imazethabenz or imazapyr.
17. The method of claim 16, wherein said imidazolinone herbicide comprises imazethapyr.
18. The method of claim 16, wherein said imidazolinone herbicide comprises imazamox.
19. The cell of any one of claims 1, 2, or 3, wherein said imidazolinone herbicide comprises at least one of: imazethapyr, imazapic, imazamox, imazaquin, imazethabenz or imazapyr.
20. The use of claim 15, wherein said imidazolinone herbicide comprises at least one of: imazethapyr, imazapic, imazamox, imazaquin, imazethabenz or imazapyr.
21. The use of claim 20, wherein said imidazolinone herbicide comprises imazethapyr.
22. The use of claim 20, wherein said imidazolinone herbicide comprises imazamox.

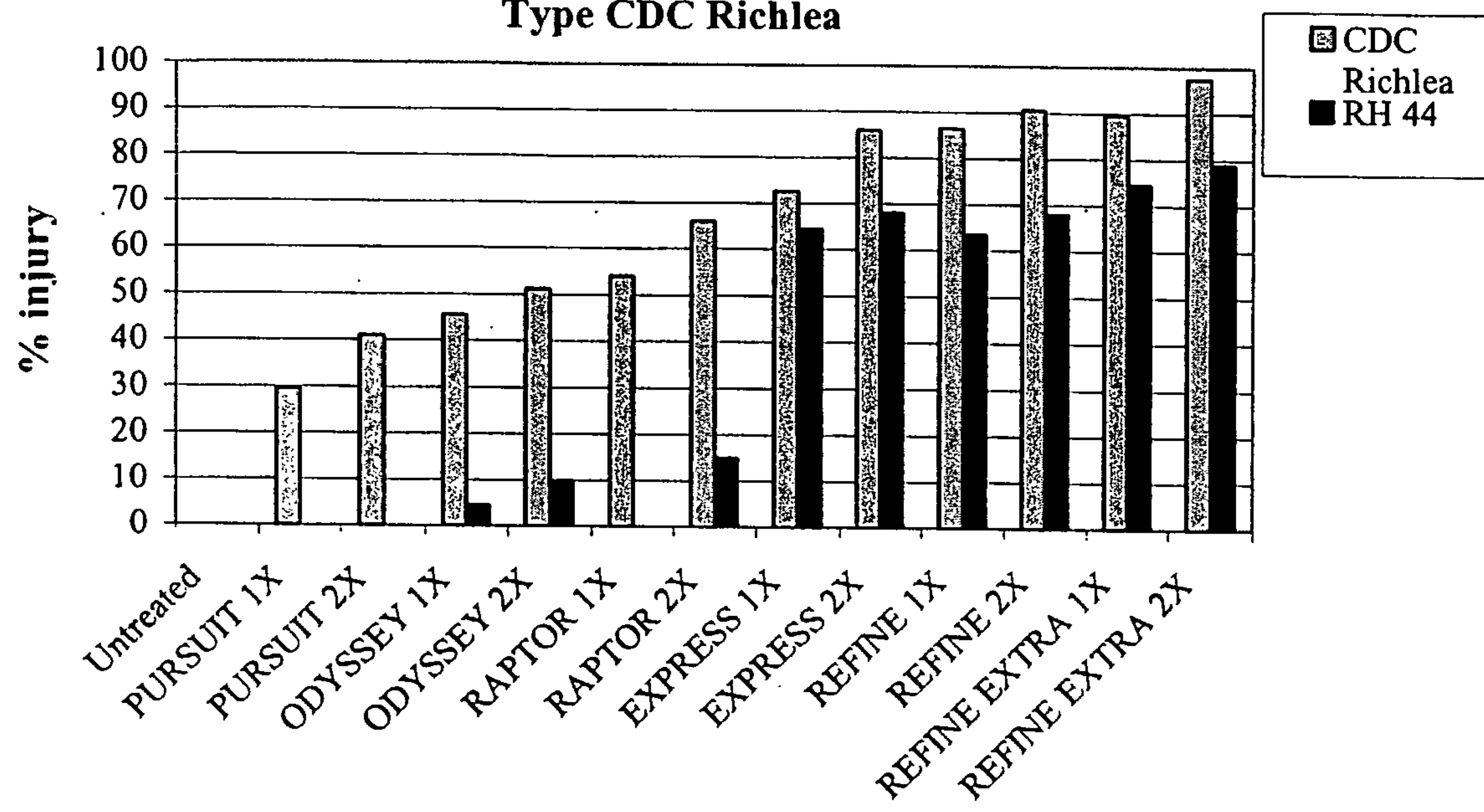
Figure 1

The change in amino acid composition and implications for sulfonylurea herbicide tolerance resulting from the gene mutation in CLEARFIELD crops conferring imidazolinone herbicide resistance

Plant	Variety	Mutation (nucleotide change)	Locus (amino acid position)	Effect on AHAS Isozyme	Tolerant to Sulfonylurea Herbicides
Corn	XI-12	Single gene point mutation AGC-AAC	621	Amino acid change serine to asparagine in AHAS isozyme	No
Canola	PM1	Single gene point mutation AGC-AAC	Unknown	Amino acid change serine to asparagine in AHAS isozyme	No
Canola	PM2	Single gene point mutation AGC-ATC	557	Amino acid change tryptophan to leucine in AHAS isozyme	Yes
Wheat	SWP965001	Single gene point mutation AGC-AAC	Gene A	Amino acid change serine to asparagine in AHAS isozyme	No

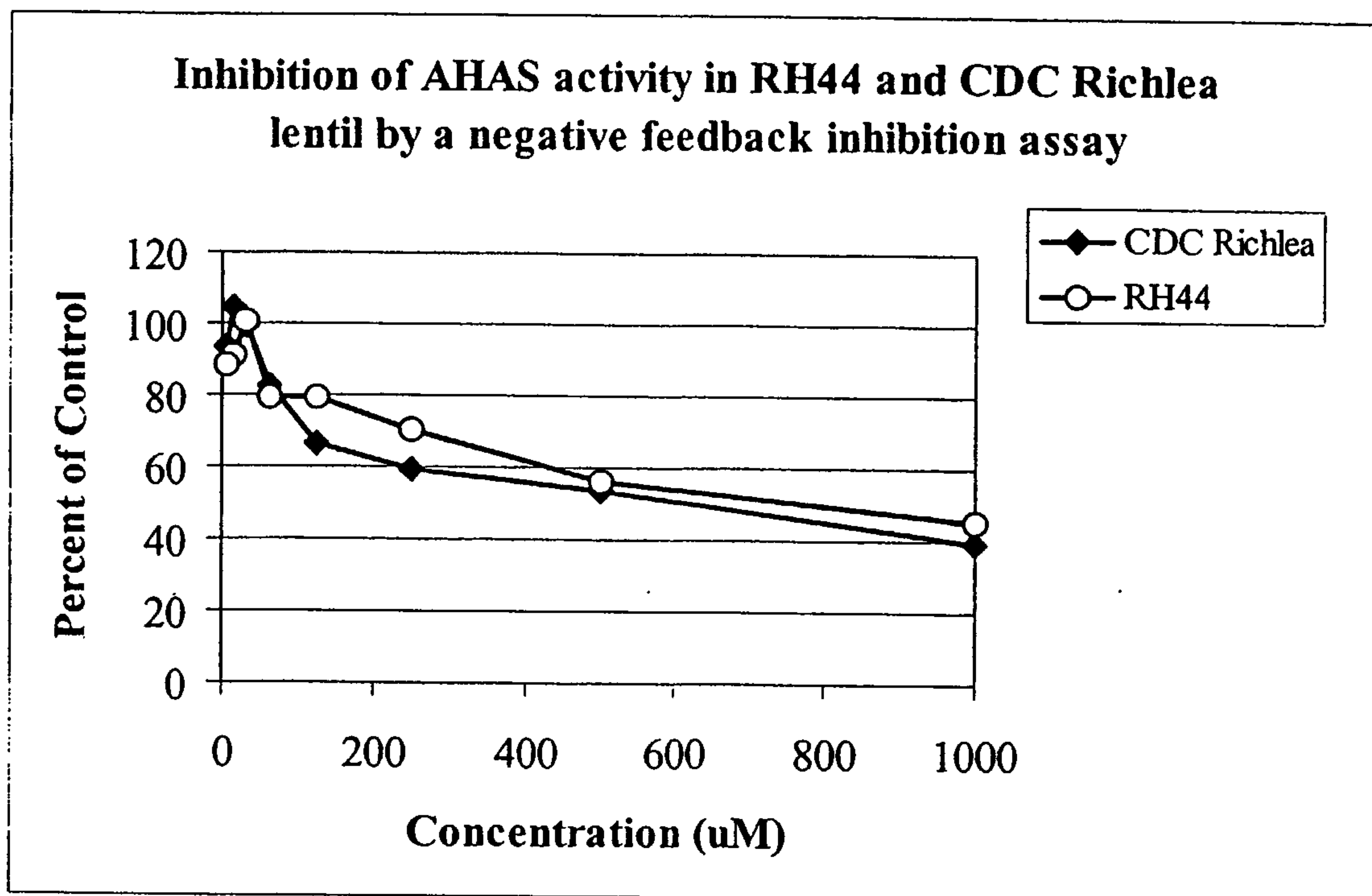
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Figure 2
Percentage of Injury of RH44 Lentil Plants as Compared to Wild
Type CDC Richlea



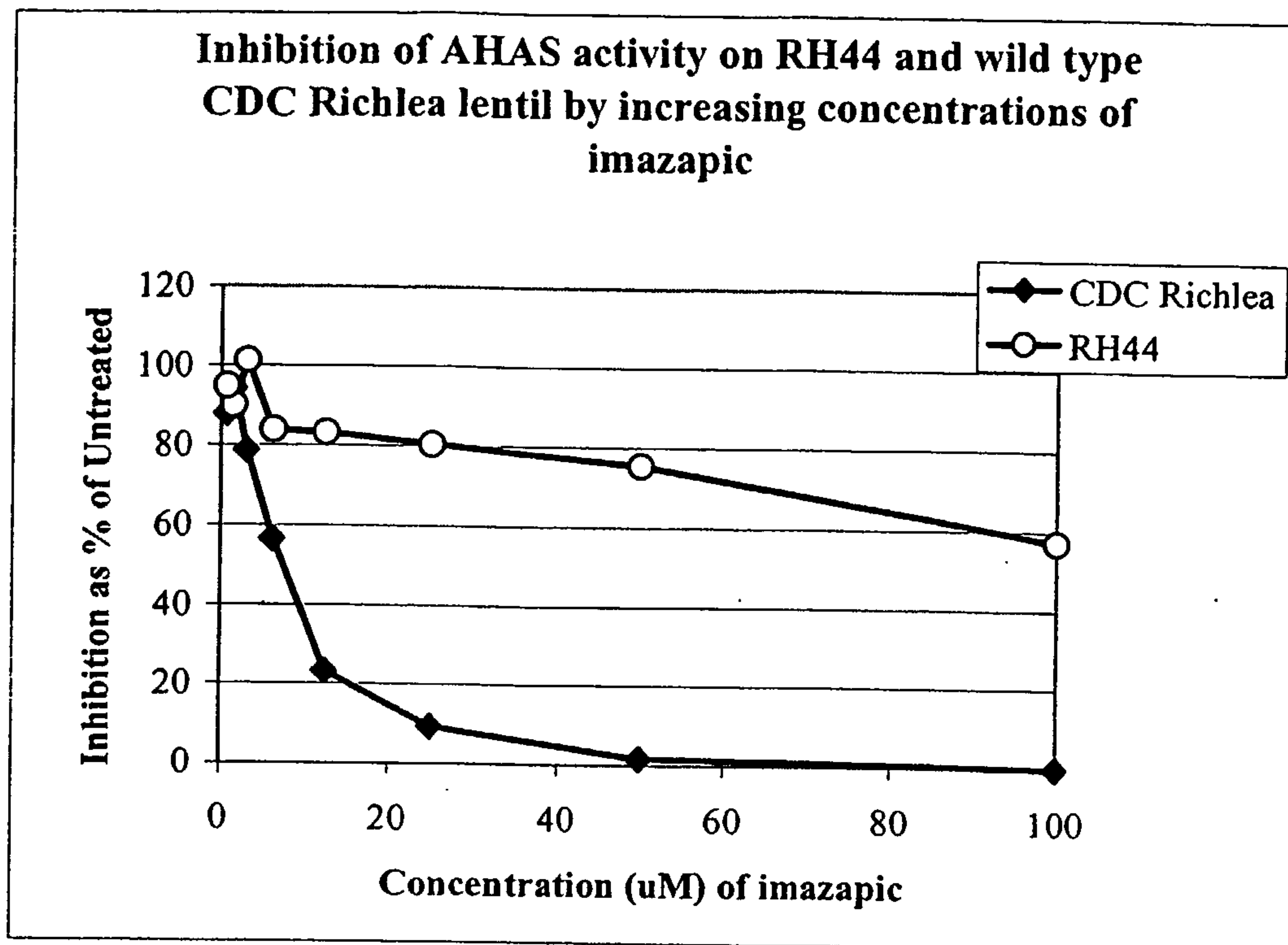
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Figure 3

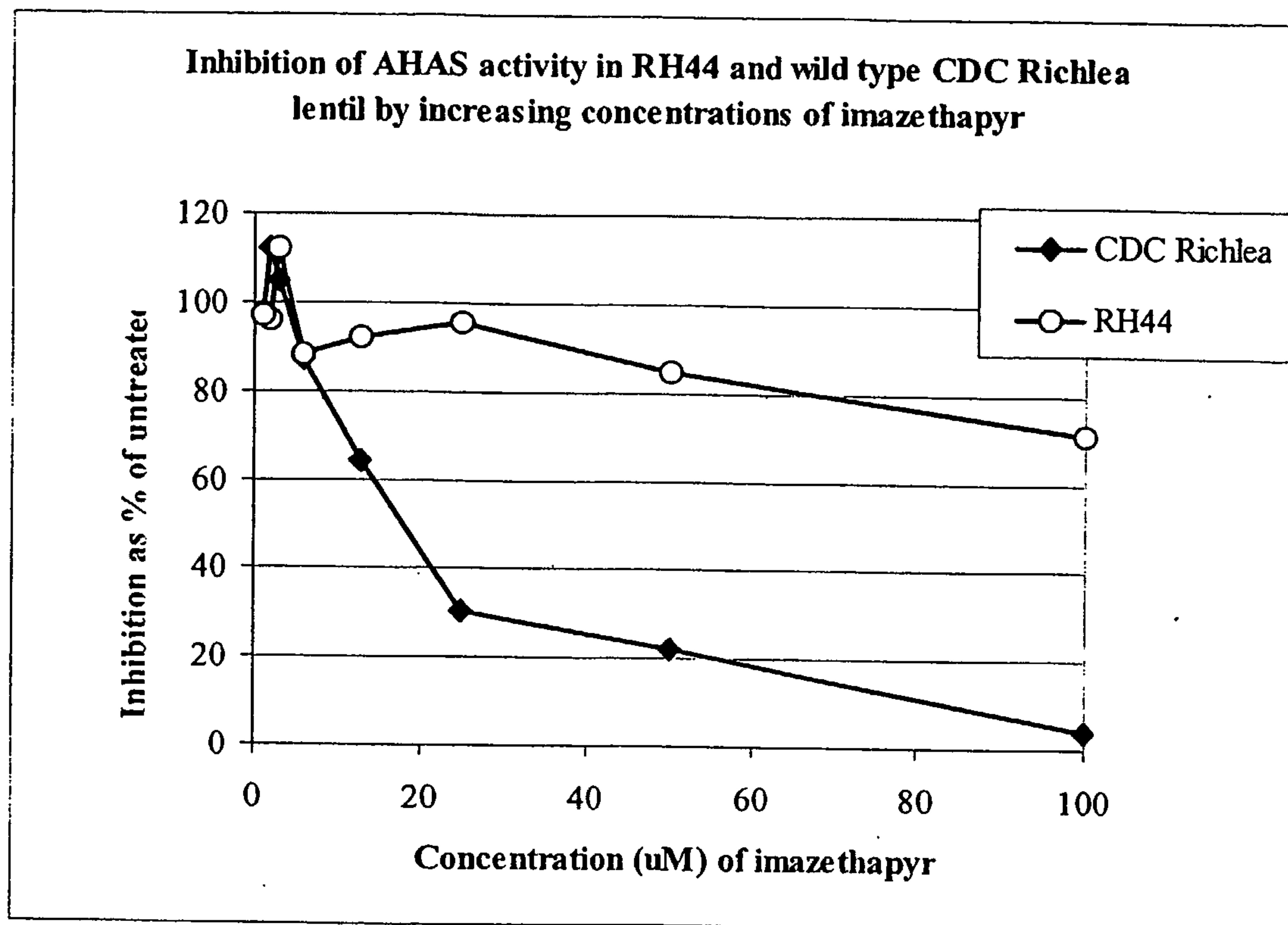


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Figure 4



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Figure 5

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Figure 6
Agronomic Traits of RH44 Lentil and Various Wild Type
Commercial Lentil Varieties

Variety	Days to flowering*	Days to maturity*	Height
CDC Sovereign	60	98	39.5
Eston	56	96(est.)	29.8
CDC Glamis	61	98	33.5
CDC Milestone	57	99	29.5
CDC Richlea	58	101	30.8
CDC Vantage	58	100	33.5
RH-44	58**	101**	33.4

* The data presented for flowering and maturity are a composite of data from several trials at various locations.

** The flowering and maturity data for RH44 are estimates based on observations in side-by-side trials with registered varieties.

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Figure 7

Effect of 2X ODYSSEY® Application on Lentil Yield
(Kernen Farm, Saskatoon, 1999 and 2000)

Line	1999 Kernen			2000 Kernen			
	Yield of control	Yield with 2X ODYSSEY	% of loss from 2X ODYSSEY application	Yield of control	Yield with 2X ODYSSEY	% of loss from 2X ODYSSEY application	% of loss from 2X ODYSSEY application
	(g/plot)	(g/plot)		(g/plot)	(g/plot)		(2 years)
CDC Richlea	1241	184	85		185	82	83
CDC Sovereign	1064	229	81		343	66	74
CDC Glamis	753	305	75		411	59	67
CDC Vantage	1114	536	57		370	63	60
Eston	1602	485	61		532	47	54
CDC Milestone	1559	936	24		500	50	37
RH-44	1345	1237	8	1000	1011	0	4
CV		29			31		

Unsprayed mean yield of RH44 lentil in year 1999 was 1345 g/plot and 1000 g/plot in year 2000. Control plots were maintained as “weed-free” by hand-weeding plots.

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Figure 8

A Comparison of the Amino Acid Composition
of RH44 Lentil and Wild Type Lentil Varieties

Test	Wild Type Lentil Varieties				RH44 Lentil				
	Average	Std Dev.	Min	Max	Sample 1	Sample 2	Sample 3	Average	Std Dev.
	%				%				
Tryptophan	0.21	0.02	0.18	0.24	0.21	0.20	0.21	0.21	0.01
Cystine	0.27	0.01	0.25	0.28	0.27	0.27	0.28	0.27	0.01
Methionine	0.20	0.01	0.19	0.21	0.20	0.20	0.20	0.20	0.00
Aspartic Acid	2.64	0.11	2.46	2.81	2.56	2.55	2.56	2.56	0.01
Threonine	0.86	0.03	0.81	0.92	0.82	0.83	0.84	0.83	0.01
Serine	1.12	0.03	1.06	1.17	1.08	1.08	1.10	1.09	0.01
Glutamic Acid	3.62	0.11	3.42	3.84	3.53	3.53	3.55	3.54	0.01
Proline	0.87	0.06	0.76	0.95	0.95	0.84	0.88	0.89	0.06
Glycine	0.93	0.03	0.88	0.97	0.93	0.91	0.93	0.92	0.01
Alanine	0.90	0.07	0.80	0.99	0.83	0.83	0.84	0.83	0.01
Valine	1.13	0.05	1.06	1.25	1.10	1.08	1.10	1.09	0.01
Isoleucine	1.00	0.03	0.95	1.05	0.96	0.95	0.97	0.96	0.01
Leucine	1.67	0.06	1.58	1.80	1.62	1.58	1.61	1.60	0.02
Tyrosine	0.45	0.02	0.40	0.48	0.44	0.43	0.45	0.44	0.01
Phenylalanine	1.12	0.05	1.04	1.24	1.08	1.08	1.08	1.08	0.00
Lysine, Total	1.56	0.04	1.49	1.63	1.51	1.50	1.53	1.51	0.02
Histidine	0.62	0.02	0.58	0.65	0.59	0.58	0.60	0.59	0.01
Arginine	1.67	0.09	1.51	1.84	1.61	1.57	1.61	1.60	0.02

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Figure 9

A Comparison of the Nutritional Composition of RH44 Lentil and Wild Type Lentil Varieties

Test	Wild Type Lentil Varieties				RH44 Lentil				
	Average	Std Dev.	Min	Max	Sample 1	Sample 2	Sample 3	Average	Std Dev.
Moisture- Forced Draft Oven	8.24	0.28	7.86	8.61	8.14	8.25	8.27	8.22	0.07
Crude Fat/Oil	0.76	0.07	0.63	0.94	0.82	0.76	0.76	0.78	0.03
Protein	25.22	0.73	23.97	26.22	25.04	23.91	24.50	24.48	0.57
Crude Fibre	4.31	0.61	3.10	5.20	4.90	4.80	4.70	4.80	0.10

Percentage of Injury of RH44 Lentil Plants as Compared to Wild Type CDC Richlea

