



# Characterization of senescence-associated protease activities involved in the efficient protein remobilization during leaf senescence of winter oilseed rape

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## ABSTRACT

Oilseed rape (*Brassica napus* L.) is a crop plant characterized by a poor nitrogen (N) use efficiency that is mainly due to low N remobilization efficiency during the sequential leaf senescence of the vegetative stage. As a high leaf N remobilization efficiency was strongly linked to a high remobilization of proteins during leaf senescence of rapeseed, our objective was to identify senescence-associated protease activities implicated in the protein degradation. To reach this goal, leaf senescence processes and protease activities were investigated in a mature leaf becoming senescent in plants subjected to ample or low nitrate supply. The characterization of protease activities was performed by using *in vitro* analysis of RuBisCO degradation with or without inhibitors of specific protease classes followed by a protease activity profiling using activity-dependent probes. As expected, the mature leaf became senescent regardless of the nitrate treatment, and nitrate limitation enhanced the senescence processes associated with an enhanced degradation of soluble proteins. The characterization of protease activities revealed that: (i) aspartic proteases and the proteasome were active during senescence regardless of nitrate supply, and (ii) the activities of serine proteases and particularly cysteine proteases (Papain-like Cys proteases and vacuolar processing enzymes) increased when protein remobilization associated with senescence was accelerated by nitrate limitation.

**Short statement:** Serine and particularly cysteine proteases (both PLCPs and VPEs) seem to play a crucial role in the efficient protein remobilization when leaf senescence of oilseed rape was accelerated by nitrate limitation.

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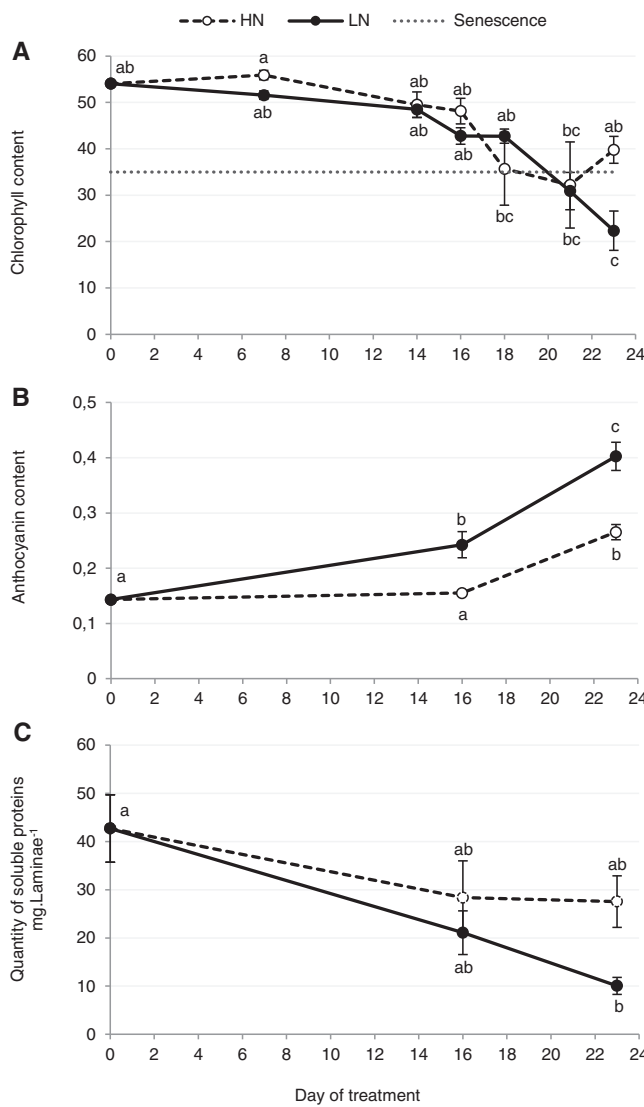
## 1. Introduction

Oilseed rape (*Brassica napus* L.) is the third largest oleaginous crop worldwide and the dominant oilseed crop in northern Europe. It is cultivated for its seeds, from which the extracted oil is used for human food and non-food uses (biofuel, detergents and lubricants) and the cake leftover from processing, which is rich in proteins and micronutrients, is used for animal feed. However, oilseed rape needs a large amount of nitrogen (N) fertilizers (160–250 kg N ha<sup>-1</sup> year<sup>-1</sup>) for its development [1], which can lead to economic losses as well as negative impacts on the environment, and so N fertilization represents the main operational cost for farmers. That is why, in a context of sustainable agriculture, a reduction in N inputs combined with the optimization of oilseed rape Nitrogen Use Efficiency (NUE) has become essential [2,3]. Oilseed rape is

**Abbreviations:** AP, aspartic protease; CCV, chloroplast vesiculation-containing vesicle; CP, cysteine protease; CXE, carboxylesterase; HN, high nitrate; LN, low nitrate; MES, methylesterase; N, nitrogen; NPC, no probe control; NRE, nitrogen remobilization efficiency; NUE, nitrogen use efficiency; PLCP, papain-like cysteine protease; POP1, prolyl oligopeptidase-like protease; RuBisCO (RBC), ribulose-1,5-biphosphate carboxylase/oxygenase; RBCL, large subunit of RuBisCO; SAP, senescence associated protease; SAV, senescence associated vesicles; SCPL, serine carboxypeptidase-like protein; VPE, vacuolar processing enzyme.

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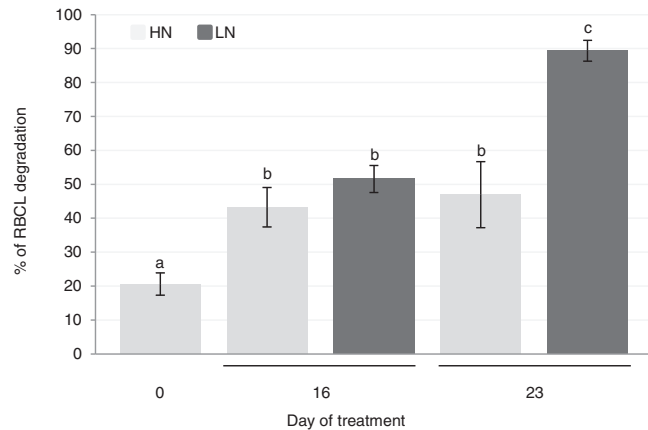
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**Fig. 1.** Changes in parameters associated with senescence in a mature leaf of *B. napus* L. supplied with ample (HN) or low (LN) nitrate for 23 days. Six-week-old plants were subjected to ample (HN: 3.75 mM  $\text{NO}_3^-$ ) or low nitrogen supply (LN: 0.375 mM  $\text{NO}_3^-$ ) for 23 days. A mature leaf that became senescent during the experiment (Leaf rank no. 12) was chosen on the basis of its leaf area determined with a LI-COR 300 area metre. Chlorophyll content was measured with a SPAD meter (A). A leaf was considered as senescent when the chlorophyll content is decreased by at least 40% of the initial value (here the senescence threshold was placed at 35 SPAD units). Anthocyanin content was measured with an optical sensor system (Multiplex®) (B) and the quantity of soluble proteins was determined after extraction by protein-dye staining [42] (C). Vertical bars indicate  $\pm$  SE of the mean ( $n=3$ ). Statistics are represented by letters ( $p<0.05$ , ANOVA, Newman–Keuls test).

characterized by a low NUE because only 50% of the N from fertilizers is finally recovered in the seeds, while a significant proportion of N inputs is returned to the environment directly or by the fallen leaves [4]. Moreover, it was shown that the low NUE is mainly due to a weak N Remobilization Efficiency (NRE) [5–7]. Indeed, during the vegetative stages of growth the recycling of foliar N is not optimal during the ‘sequential’ leaf senescence, which corresponds to a senescence progression along the axis of the plant that affects leaves as they reach maturity and leads to nutrient remobilization from the older leaves to the younger leaves [3].

In plants, senescence corresponds to the final stage of leaf development and is characterized by the transition from assimilation to remobilization of nutrients [8]. This phenomenon contributes to resource management, recycling and nutrient remobilization

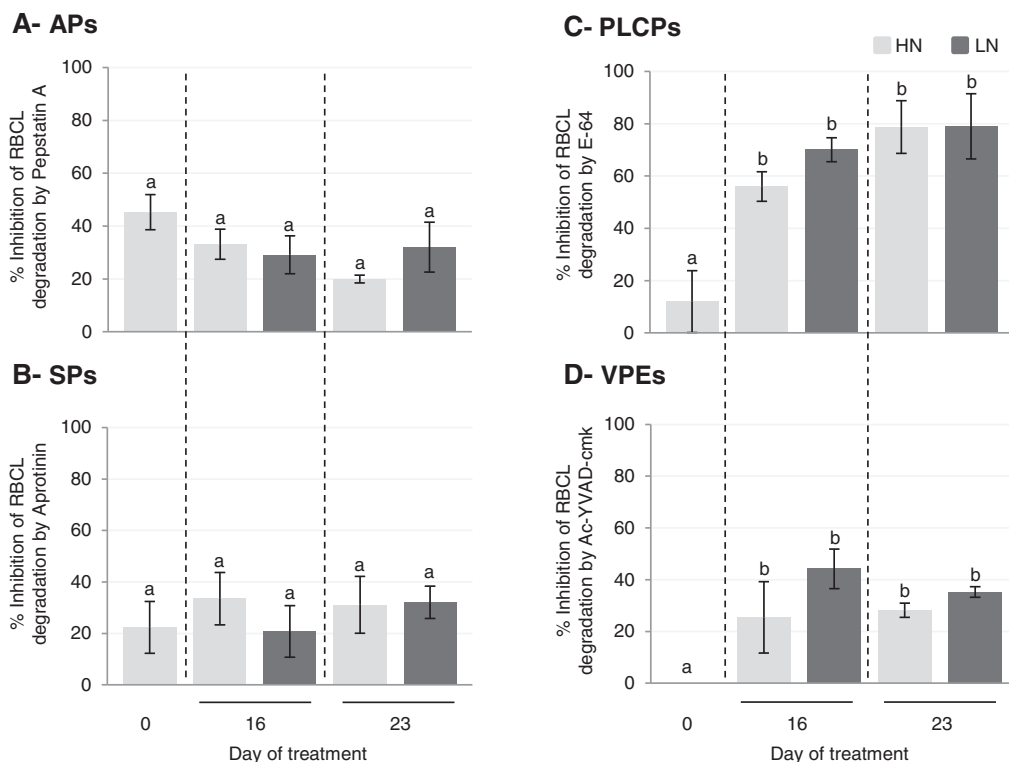


**Fig. 2.** The degradation of RuBisCO (RBCL) at pH 5.5 increases during leaf senescence of *B. napus* L. subjected to N limitation (LN) compared to plants with ample nitrate supply (HN). Soluble proteins were extracted after 0, 16 and 23 days of HN (3.75 mM  $\text{NO}_3^-$ ) or LN (0.375 mM  $\text{NO}_3^-$ ) treatment and were incubated in sodium acetate buffer (50 mM, pH 5.5) for 30 min at 37 °C. The reaction was stopped by adding ice-cold acetone. To determine the initial quantity of RuBisCO (RBCL, large subunit) another sample was also treated as described above but the reaction was stopped immediately. Samples were separated on SDS-PAGE Stain-free gels (Mini-PROTEAN® TGXTM Stain Free), scanned under UV light with a Gel Doc™ EZ scanner and analysed (for details see Section 2). The degradation of RBCL was calculated as the difference in the quantity between non incubated and incubated samples and expressed as % of RBCL degradation. Vertical bars indicate  $\pm$  SE of the mean ( $n=3$ ). Statistics are represented by letters ( $p<0.05$ , ANOVA, Newman–Keuls test).

efficiency [9] and is essential for plant productivity [10,11]. Leaf senescence, controlled by intrinsic and environmental factors, leads to a sequence of events such as chlorophyll loss, degradation of macromolecules like proteins, dismantling of cellular components, and cell death [3,12,13]. Considered as the most important degradation process during leaf senescence, protein breakdown allows the remobilization of N [14] and the resulting amino acids or peptides are exported to growing parts of the plant via the phloem, leading to an increase in the concentration of amino acids in the phloem sap [15,16].

A recent study has shown that the enhancement of amino acid export and soluble protein degradation in senescing leaves of oilseed rape are crucial for the improvement of N remobilization [17]. This study on the genotypic variability of foliar N remobilization at the vegetative stage of *B. napus* L. revealed that the export of amino acids is efficient for the ten genotypes studied. Otherwise, the genotype Aviso, unlike other genotypes, is able to maintain its leaf biomass production in response to low nitrate supply and this was essentially due to an improvement of soluble protein degradation. Up to 75% of leaf N is located in chloroplasts as proteins, especially in RuBisCO (ribulose-1,5-biphosphate carboxylase/oxygenase, EC 4.1.1.39) and LHCII (Light Harvesting complex II) [18,19] located in the stroma or in the thylakoid membrane, respectively. Moreover, it is known that the RuBisCO degradation during senescence provides much of N needed to the development of growing organs [20,21]. Accordingly, the improvement of RuBisCO recycling by proteases is crucial for the optimization of N remobilization [19]. Surprisingly, in fallen leaves of *B. napus* L., RuBisCO corresponds to one of the major residual proteins [22], suggesting that proteolysis is limiting for N remobilization during sequential senescence in leaves of oilseed rape.

There are many protease classes involved in protein breakdown during senescence including serine, aspartic, metallo- and cysteine proteases and the proteasome [23]. In wheat, several serine proteases are induced in response to N starvation during leaf senescence [24]. Moreover, N starvation in oilseed rape plants leads to an increase in an aspartic protease during the first phases of leaf senes-



**Fig. 3.** Changes in activities of aspartic proteases (APs, A), serine proteases (SPs, B), papain-like Cys proteases (PLCPs, C) and vacuolar processing enzymes (VPEs, D) at pH 5.5 during leaf senescence of *B. napus* L. supplied with ample (HN) or low (LN) nitrate for 23 days. Soluble proteins were extracted after 0, 16 and 23 days of HN (3.75 mM  $\text{NO}_3^-$ ) or LN (0.375 mM  $\text{NO}_3^-$ ) treatment and were incubated in sodium acetate buffer (50 mM, pH 5.5) for 30 min at 37 °C without inhibitor or in the presence of 50  $\mu\text{M}$  of Pepstatin A (inhibitor of aspartic proteases), Aprotinin (inhibitor of serine proteases), E-64 (specific inhibitor of PLCPs) or Ac-YVAD-cmk (inhibitor of VPEs). The reaction was stopped by adding ice-cold acetone. To determine the initial quantity of RuBisCo (large subunit, RBCL), another sample was also treated as described above but the reaction was stopped immediately. Samples were separated on SDS-PAGE Stain-free gels (Mini-PROTEAN® TGXTM Stain Free), scanned under UV light with a Gel Doc™ EZ scanner and analysed (for details see Section 2). The degradation of RBCL was calculated as the difference in quantity between non incubated and incubated samples and expressed as % of RBCL degradation, while the % of inhibition due to the different inhibitors was calculated as the difference in degradation in the presence or absence of inhibitors. Vertical bars indicate  $\pm$  SE of the mean ( $n=3$ ). Statistics are represented by letters ( $p<0.05$ , ANOVA, Newman-Keuls test).

cence [25]. Furthermore, aspartic protease CND41 participates in RuBisCO degradation during senescence and *CND41* antisense tobacco presented a delay in the senescence process and an accumulation of N in senescent leaves [26,27]. Metalloproteases have also been implicated in senescence and several genes encoding metalloproteases, particularly FtsH proteases, are induced in senescent leaves of *Arabidopsis thaliana* [28]. Using proteomics, Desclos et al. [25] showed that nitrate limitation induces a chloroplastic FtsH in senescent leaves of oilseed rape (cv. Capitol). Cysteine proteases might be crucial for the degradation of proteins as they are the most abundant class of proteases up-regulated during leaf senescence-related proteolysis [28,29]. Indeed, in senescent leaves of *A. thaliana*, a high cysteine protease activity was detected in Senescence Associated Vesicles (SAV) [30] and several experiments have demonstrated a role for this class of proteases in RuBisCO degradation [21,31]. Some studies implicated Vacuolar Processing Enzymes (VPEs, a sub-family of cysteine proteases) in leaf senescence. As demonstrated by Sanmartín et al. [32],  $\alpha$ VPEs and  $\gamma$ VPEs were up-regulated in senescing vegetative organs of *A. thaliana* and encoded proteins might be responsible for the activation of downstream proteases involved in the recycling of amino acids during senescence [33]. Finally, the ubiquitin-proteasome system is also implicated in protein degradation during senescence. The proteasome is able to degrade ubiquitinated, short-lived, regulatory or abnormal proteins. The proteasome consists of the 19S regulatory particle and the 20S core protease. The 20S core protease contains the catalytic subunits  $\beta 1$  (caspase-like activity),  $\beta 2$  (trypsin-like activity) and  $\beta 5$  (chymotrypsin-like activity) [34]. Proteomic anal-

ysis revealed that the catalytic  $\beta 1$  subunit was induced during leaf senescence in oilseed rape [3,25].

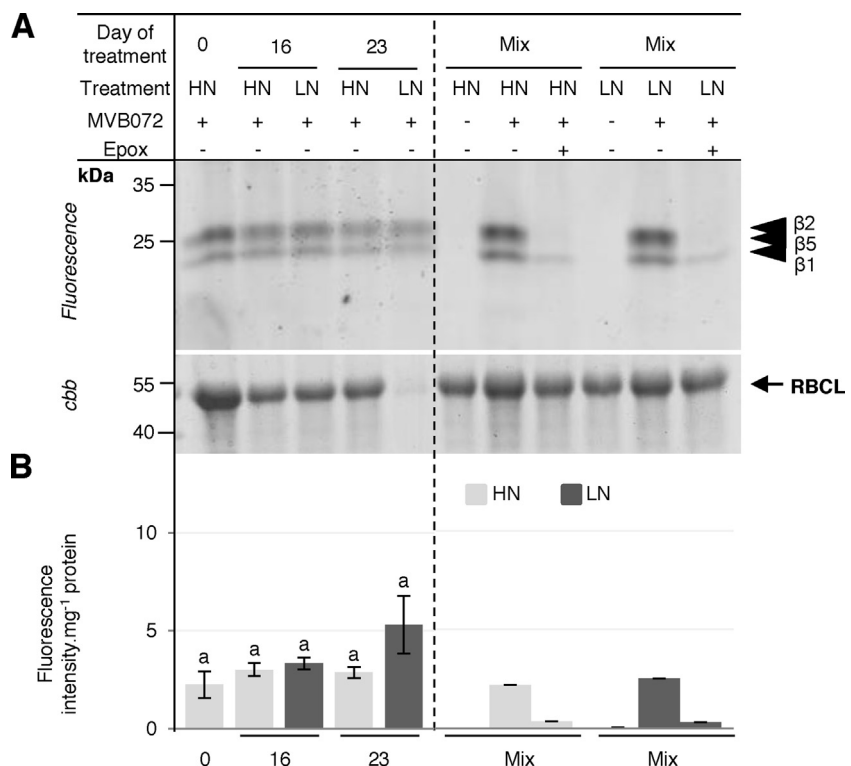
While RuBisCO degradation during senescence is relatively well studied in other plant species [19,26,35], this process remains largely unknown in oilseed rape. Because soluble protein degradation in senescing leaves of oilseed rape is crucial for the improvement of leaf N remobilization, the characterization of protease activities is key for comprehension of N remobilization. Thus, our objective was to identify senescence-associated protease activities implicated in protein remobilization during leaf senescence at the vegetative stage.

As proteases are tightly regulated to prevent damage by uncontrolled proteolytic activities, it is difficult to predict the activity of proteases on the basis of their transcript or protein abundance alone. That is why we focused this work on detecting protease activities by performing an *in vitro* analysis of the degradation of the RuBisCO large subunit (RBCL), with or without inhibitors of specific protease classes. Additionally, to identify the active proteases associated with the efficient leaf protein degradation during leaf senescence, we used activity-based protein profiling, an original method which allows the detection and identification of proteases in their active states [36].

## 2. Material and methods

### 2.1. Chemicals

E-64, epoxomycin, Ac-YVAD-cmk, diisopropylfluorophosphate (DFP), aprotinin and pepstatin A were from SIGMA-ALDRICH®. The



**Fig. 4.** Proteasome activity at pH 7.5 is stable during leaf senescence of *B. napus* L. supplied with low nitrate (LN) concentration for 23 days. Soluble proteins were extracted after 0, 16 and 23 days of HN (3.75 mM NO<sub>3</sub><sup>-</sup>) or LN (0.375 mM NO<sub>3</sub><sup>-</sup>) treatment and were subjected to standard protease activity profiling with MVB072 (specific fluorescent probe of proteasome) (pH 7.5; 1 h labelling). These extracts were separated by SDS-PAGE and scanned to detect fluorescence (A). The Coomassie brilliant blue-stained (cbb) protein gel shows the total amount of input proteins after incubation. Mix corresponds to a mix of 0, 16 and 23 days extracts (HN or LN) in the presence of MVB072 alone, MVB072 and epoxomicin (specific inhibitor of proteasome) or in the absence of MVB072 (no probe control). The fluorescence intensity representative of the proteasome activity was calculated relative to the protein amount (B). Black arrowheads correspond to the position of catalytic subunits β1, β2 and β5 of the proteasome. The gel presented in this figure is representative of three biological replicates. Vertical bars indicate ± SE of the mean of three biological replicates. Statistics are represented by letters (*p* < 0.05, ANOVA, Newman–Keuls test).

**Table 1**  
Different classes of proteases studied in this work, their specific inhibitors and activity-based probes. The codification of the probes and the corresponding references for the synthesis of these probes are listed.

Class of protease	Sub-family	Inhibitor	Probe with fluorescent tag	Probe with biotin tag	References for the probes
Cys proteases	PLCPs (C1)	E-64	MV201	DCG04	[37]
	VPE (C14)	Ac-YVAD-cmk	JOPD1	–	[38]
Ser proteases	Subtilisins (S8), POPPs (S9), SCPLs (S10)	DFP or Aprotinin	FP-Rh	FP-biotin	[39]
Proteasome	T3	Epoxomicin	MVB072	–	[40]
Asp proteases	–	Pepstatin A	–	–	–

probes MV201, JOPD1, MVB072, FP-Rh, DCG-04 and FP-biotin were available in the laboratory and described in Table 1 [37–40].

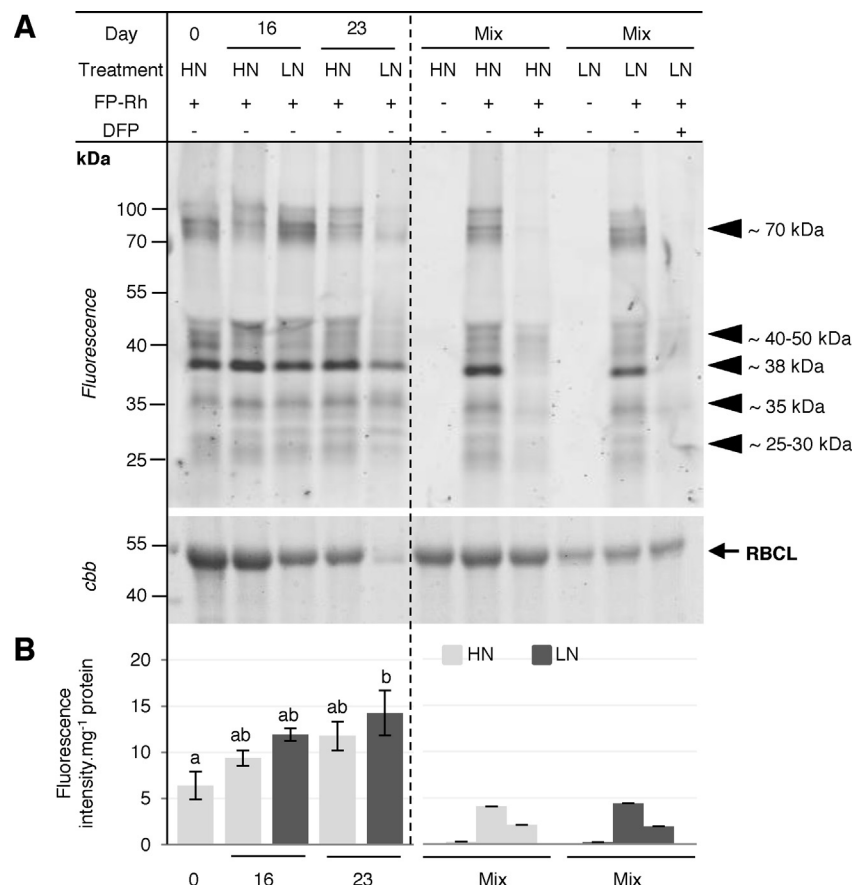
## 2.2. Plant material and growth conditions

Oilseed rape (*B. napus* L. genotype Aviso) plants were cultivated at the vegetative stages in a greenhouse under a 16 h light regime at 20 °C (day)/15 °C (night) with a PAR (Photosynthetically Active Radiation) of 400 μmoles photon. s<sup>-1</sup> m<sup>-2</sup> at the canopy. After germination, seedlings were transferred into 2.5 L pots containing mixed vermiculite/perlite (1:2 v/v) and cultivated with 25% Hoagland nutrient solution (1.25 mM Ca(NO<sub>3</sub>)<sub>2</sub>·4H<sub>2</sub>O, 1.25 mM KNO<sub>3</sub>, 0.5 mM MgSO<sub>4</sub>, 0.25 mM KH<sub>2</sub>PO<sub>4</sub>, 0.2 mM EDTA·2NaFe·3H<sub>2</sub>O, 14 μM H<sub>3</sub>BO<sub>3</sub>, 5 μM MnSO<sub>4</sub>, 3 μM ZnSO<sub>4</sub>, 0.7 μM (NH<sub>4</sub>)<sub>6</sub>Mo<sub>7</sub>O<sub>24</sub>, 0.7 μM CuSO<sub>4</sub>, 0.1CoCl<sub>2</sub>). After six weeks (Day 0 (D0) of the experiment), plants were supplied with 25% Hoagland solution containing two different concentrations of nitrate: high (HN: 3.75 mM) or low nitrate (LN: 0.375 mM NO<sub>3</sub><sup>-</sup> with a compensation of Ca and K elements by adding 1.25 mM CaCl<sub>2</sub>·2H<sub>2</sub>O and 0.875 mM KCl). Leaves were numbered in order

of their appearance where leaf rank no. 1 (L1) was the oldest leaf. At D0, a mature leaf becoming senescent during the experiment (leaf rank no. 12 (L12)) was chosen on the basis of its leaf area determined with a LI-COR 300 area meter (LI-COR, Lincoln, NE, USA) and chlorophyll content with a SPAD meter (Soil Plant Analysis Development; Minolta, SPAD-502 model): mean leaf area value of (62.49 cm<sup>2</sup> ± 9.16% variation; mean SPAD value of 52.20 ± 5.52% variation). Plants were harvested after 0, 16 and 23 days of treatment (D0, D16 and D23) and anthocyanin levels of L12 were measured before each harvest by an optical sensor system (Multiplex®, Orsay, France) as previously described by D'Hoooge et al. [41]. Laminae of L12 were frozen (−80 °C) and used for proteomics and molecular analyses.

## 2.3. Protease activity associated with degradation of RuBisCO (RBCL)

Soluble proteins were extracted by grinding 200 mg of frozen leaf tissue with 500 μL citrate-phosphate buffer (20 mM citrate, 160 mM phosphate, pH 6.8 containing 50 mg



**Fig. 5.** Activity of serine proteases at pH 7.5 is up-regulated during leaf senescence of *B. napus* L. supplied with low (LN) nitrate for 23 days. Soluble proteins were extracted after 0, 16 and 23 days of HN ( $3.75 \text{ mM } \text{NO}_3^-$ ) or LN ( $0.375 \text{ mM } \text{NO}_3^-$ ) treatment and were subjected to standard protease activity profiling with FP-Rh (specific fluorescent probe of serine proteases) (pH 7.5; 1 h labelling). These extracts were separated by SDS-PAGE and scanned to detect fluorescence (A). The Coomassie brilliant blue-stained (cbb) protein gel shows the total amount of input proteins after incubation. The fluorescence intensity representative of serine protease global activity was calculated relative to the protein amount (B). Mix corresponds to a mix of 0, 16 and 23 days extracts (HN or LN) in the presence of FP-Rh alone, FP-Rh and DFP (specific inhibitor of serine proteases) or in the absence of FP-Rh (no probe control). Black arrowheads correspond to proteases already active at day 0. The gel presented in this figure is representative of three biological replicates. Vertical bars indicate  $\pm \text{SE}$  of the mean of three biological replicates. Statistics are presented by letters ( $p < 0.05$ , ANOVA, Newman-Keuls test).

of polyvinylpolypyrrolidone (PVPP)). After centrifugation (1 h,  $12,000 \times g$ ,  $4^\circ\text{C}$ ), the resulting supernatant containing proteins was used for determination of the quantity of soluble proteins by protein-dye staining [42] using bovine serum albumin (BSA) as standard.

To identify the proteases classes associated with protein remobilization during leaf senescence of plants subjected to nitrate limitation (LN) or ample nitrate supply (HN), the degradation of RuBisCo large subunit (RBCL) by proteases within the soluble protein extract was studied with or without inhibitors of specific protease classes by using a method modified from Girondé et al. [43] (Supplemental data; Fig. S1). In this method, the RBCL was used as a target of proteolysis because this stromal soluble protein is one of the main substrate of proteases during leaf senescence. Protease activities were determined at pH 5.5 and 7.5. To achieve this, 8 or 12  $\mu\text{g}$  of proteins were incubated in a 200  $\mu\text{L}$  final volume with sodium acetate buffer (50 mM, pH 5.5) in the presence or absence of 50  $\mu\text{M}$  of E-64, Ac-YVAD-cmk, aprotinin or pepstatin A (Table 1). Otherwise, to study cysteine proteases, 2 mM of dithiothreitol (DTT) were added in this mix. The incubation was performed for 30 min at  $37^\circ\text{C}$  under gentle agitation. Alternatively, 8 or 12  $\mu\text{g}$  of proteins were incubated for 90 min at  $37^\circ\text{C}$  in a 200  $\mu\text{L}$  final volume with Tris-base buffer (125 mM; pH 7.5) without inhibitor or in the presence of 50  $\mu\text{M}$  of epoxomicin. An equal volume of dimethylsulfoxide (DMSO) was added to the No-Inhibitor-Control. Degradation was stopped by adding

1 mL of ice-cold acetone. The pellet obtained after centrifugation (15 min,  $16,000 \times g$ ,  $4^\circ\text{C}$ ) was dissolved in 2X SDS-PAGE gel loading buffer (140 mM sodium dodecyl sulfate, 200 mM Tris, 20% glycerol, 5%  $\beta$ -mercaptoethanol, 0.3 mM Bromophenol Blue) and heated at  $90^\circ\text{C}$  for 10 min. To determine the initial quantity of RBCL, the protein extract was also treated without inhibitor and the proteolytic reaction was stopped immediately by adding 1 mL of ice-cold acetone, as described above (supplemental Fig. S1). Soluble protein extracts were separated on a 4–15% gradient in SDS-PAGE precast Stain-free gels (Mini-PROTEAN<sup>®</sup> TGX<sup>TM</sup> Stain Free, Bio-Rad, Marne-la-Coquette, France) and scanned under UV light with a Gel Doc<sup>™</sup> EZ scanner (Bio-Rad, Marne-la-Coquette, France). The amount of RBCL (expressed as volume) was quantified by using ImageLab<sup>™</sup> software (Bio-Rad, Marne-la-Coquette, France) according to the manufacturer's instructions. The percentage of RBCL degradation was calculated as the difference in quantity between non incubated and incubated samples without inhibitors. The percentage of inhibition due to the different inhibitors was calculated as the difference in degradation without or in the presence of inhibitors.

#### 2.4. Protease activity profiling of extracts

Soluble proteins were extracted by grinding 200 mg of frozen leaf tissue in a microtube and mixing with 1 mL of water. The extracts were cleared by centrifugation (5 min,  $13,000 \times g$ ,  $4^\circ\text{C}$ ).

**Table 2**LC-MS/MS identification of serine proteases labelled with the probe FP-biotin in a senescent leaves of *B. napus* L. after 23 days of nitrate limitation.

Cutting zone (kDa)	Protein accession no. [ <i>Brassica napus</i> ]/NCBI accession no.	Score	Matches	Match (sig)	Exp. Mass (Da)	Theo. Mass (Da)	Result of BLAST protein [ <i>Brassica napus</i> ]-protein [organism]/NCBI accession no./% identity	Class
<b>70</b>	<b>Serinepeptidases</b>							
	BnaC05g15670D [ <i>Brassica napus</i> ]/gi 674888482	91	5	4	82234	81657.74	PREDICTED: subtilisin-like protease [ <i>Brassica rapa</i> ]/gi 685318103/99% Serine-type endopeptidase SBT5.2 [ <i>Arabidopsis thaliana</i> ]/gi 18394832/88%	Subtilisins S8
	BnaA09g07410D [ <i>Brassica napus</i> ]/gi 674914517	50	8	4	79767	79246.24	PREDICTED: subtilisin-like protease [ <i>Brassica rapa</i> ]/gi 685354526/85% Subtilisin-like protease [ <i>Arabidopsis thaliana</i> ]/gi 18425181/80%	Subtilisins S8
	BnaCnng55020D [ <i>Brassica napus</i> ]/gi 674865926	41	6	3	79945	79310.97	PREDICTED: subtilisin-like protease [ <i>Brassica rapa</i> ]/gi 685354526/88% Subtilisin-like protease [ <i>Arabidopsis thaliana</i> ]/gi 18425181/80%	Subtilisins S8
	BnaA03g44620D [ <i>Brassica napus</i> ]/gi 674934691	37	4	2	145873	144766.55	PREDICTED: LOW QUALITY PROTEIN: tripeptidyl-peptidase 2-like [ <i>Brassica rapa</i> ]/gi 685292411/97% Tripeptidyl peptidase II [ <i>Arabidopsis thaliana</i> ]/gi 30685230/92%	Subtilisins S8
<b>70</b>	<b>Serinepeptidases</b>							
	BnaC08g09650D [ <i>Brassica napus</i> ]/gi 674896006	193	14	6	91150	90237.22	PREDICTED: acylamino-acid-releasing enzyme-like isoform X1 [ <i>Brassica rapa</i> ]/gi 685342486/95% Acylamino acid-releasing enzyme [ <i>Arabidopsis thaliana</i> ]/gi 42566792/84%	POPLs S9
	BnaA08g30180D [ <i>Brassica napus</i> ]/gi 674878293	105	9	4	90670	89814.83	PREDICTED: acylamino-acid-releasing enzyme-like isoform X1 [ <i>Brassica rapa</i> ]/gi 685342486/98% Acylamino acid-releasing enzyme [ <i>Arabidopsis thaliana</i> ]/gi 42566792/84%	POPLs S9
	BnaC06g11680D [ <i>Brassica napus</i> ]/gi 674926851	74	15	8	91270	90528.61	PREDICTED: acylamino-acid-releasing enzyme-like isoform X2 [ <i>Brassica rapa</i> ]/gi 685307956/98% Acylamino acid-releasing enzyme [ <i>Arabidopsis thaliana</i> ]/gi 42566792/84%	POPLs S9
	BnaCnng28400D [ <i>Brassica napus</i> ]/gi 674876980	73	13	4	82654	81850.38	PREDICTED: prolyl endopeptidase-like [ <i>Brassica rapa</i> ]/gi 685274241/97% Prolyl oligopeptidase [ <i>Arabidopsis thaliana</i> ]/gi 79382269/90%	POPLs S9
	BnaA06g02850D [ <i>Brassica napus</i> ]/gi 674910247	52	5	1	81086	80567.57	PREDICTED: prolyl endopeptidase-like [ <i>Brassica rapa</i> ]/gi 685314489/99% Prolyl oligopeptidase family protein [ <i>Arabidopsis thaliana</i> ]/gi 18403046/92%	POPLs S9
	BnaA09g04940D [ <i>Brassica napus</i> ]/gi 674913439	49	2	2	82828	82423.77	PREDICTED: dipeptidyl peptidase 8 [ <i>Brassica rapa</i> ]/gi 685260765/99% Prolyl oligopeptidase family protein [ <i>Arabidopsis thaliana</i> ]/gi 15237923/87%	POPLs S9

Table 2 (Continued)

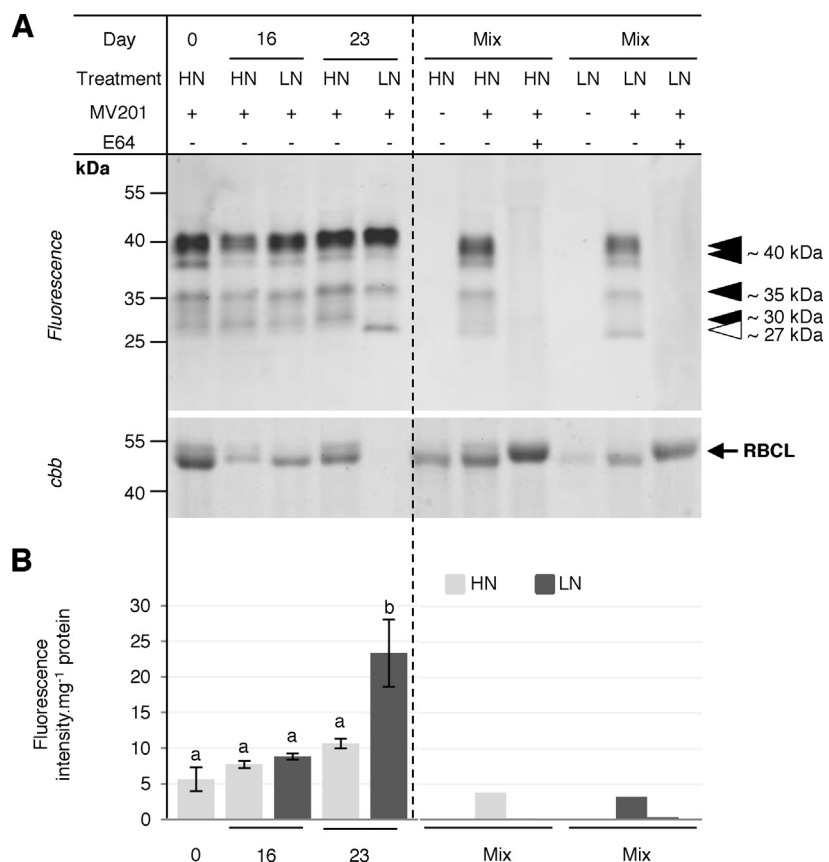
Cutting zone (kDa)	Protein accession no. [Brassica napus]/NCBI accession no.	Score	Matches	Match (sig)	Exp. Mass (Da)	Theo. Mass (Da)	Result of BLAST protein [Brassica napus]-protein [organism]/NCBI accession no./% identity	Class
	BnaA07g32380D [Brassica napus]/gi 674946140	34	9	1	82658	81854.18	PREDICTED: prolyl endopeptidase [Brassica rapa]/gi 685338483/99% Prolyl oligopeptidase [Arabidopsis thaliana]/gi 79382269/93%	POPLs S9
70	<b>Other serine hydrolases</b>							
	BnaA06g19010D [Brassica napus]/gi 674894935	64	9	6	35699	35550.37	PREDICTED: probable carboxylesterase 12 [Brassica rapa]/gi 685320048/99% Probable carboxylesterase 12 [Arabidopsis thaliana]/gi 15228425/80%	CXE
	BnaC07g46400D [Brassica napus]/gi 674961158	53	3	3	29121	31685.13	PREDICTED: LOW QUALITY PROTEIN: methylesterase 9-like [Brassica rapa]/gi 685295161/93% Methylesterase 9 [Arabidopsis thaliana]/gi 15235445/81%	MES
	BnaC01g01910D [Brassica napus]/gi 674964800	35	2	2	28978	31540.79	PREDICTED: methylesterase 9-like [Brassica rapa]/gi 685256421/95% Methylesterase 9 [Arabidopsis thaliana]/gi 15235445/76%	MES
35–40	BnaA06g19010D [Brassica napus]/gi 674894935	56	13	1	35699	35550.37	PREDICTED: probable carboxylesterase 12 [Brassica rapa]/gi 685320048/99% Probable carboxylesterase 12 [Arabidopsis thaliana]/gi 15228425/80%	CXE
	BnaA02g33270D [Brassica napus]/gi 674902269	52	3	2	17528	17424.86	PREDICTED: probable carboxylesterase 120 [Brassica rapa]/gi 685278524/100% Carboxyesterase 20 [Arabidopsis thaliana]/gi 15241725/63%	CXE
	BnaC07g22410D [Brassica napus]/gi 674930271	44	4	2	38711	38449.32	PREDICTED: probable carboxylesterase 7 isoform X1 [Brassica rapa]/gi 685325292/88% Probable carboxylesterase 7 [Arabidopsis thaliana]/gi 15227669/70%	CXE
	BnaC07g46400D [Brassica napus]/gi 674961158	45	3	2	29121	31685.13	PREDICTED: LOW QUALITY PROTEIN: methylesterase 9-like [Brassica rapa]/gi 68529516/93% Methylesterase 9 [Arabidopsis thaliana]/gi 15235445/81%	MES
	BnaA01g00900D [Brassica napus]/gi 674958220	43	5	2	28956	28632.94	PREDICTED: methylesterase 9-like [Brassica rapa]/gi 685256421/99% Methylesterase 9 [Arabidopsis thaliana]/gi 15235445/77%	MES

To characterize the serine proteases observed by fluorescent bands on gels (Fig. 5), a pull-down of biotinylated proteins was prepared after labelling with biotin-tagged probe FP and detected zones were excised, treated and analysed by LC-MS/MS (for details see Section 2). LC-MS/MS spectra were searched against the *Brassica napus* L. database using MASCOT 2.5.0 and only scores higher or equal to 30 were considered. The assigned protein of best match is provided with the GenBank accession number. Score, peptide matches, significant peptide matches (Match(sig)), experimental mass and theoretical mass are also presented. Protein sequences were also matched against sequences of *Brassicaceae* proteins using the NCBI Blast Protein Database (algorithm blastp) and better results of blasts were presented with the name of the protein, the organism, the gene accession number and the percentage of sequence identity. Finally, proteins were classified according to MEROPS database.

The concentration of soluble proteins extract in equivalent serum albumin bovine (BSA) was quantified by protein-dye staining [42].

The *in vitro* labelling of protease activities was carried out by incubating 20 µL of protein extract in a 200 µL final volume mix containing 0.5 µM of probe (MV201 or JOPD1), 50 mM of NaAc buffer (pH 5.5) and 2 mM DTT. Alternatively, 20 µL of protein extract were incubated in a 200 µL final volume of 50 mM Tris-base

buffer (pH 7.5) containing 0.5 µM of MVB072 or 0.25 µM of FP-Rh. Samples were incubated for 4 h (MV201 or JOPD1) or 1 h (MVB072 or FP-Rh) in the dark under gentle agitation. A mixture of equal volumes of soluble protein extracts of leaves (D0, D16 and D23) treated under HN or LN conditions was used as control, and 20 µL of each mixture were treated as described above. An equal volume of DMSO was added to the No-Probe-Control (NPC). Moreover, com-

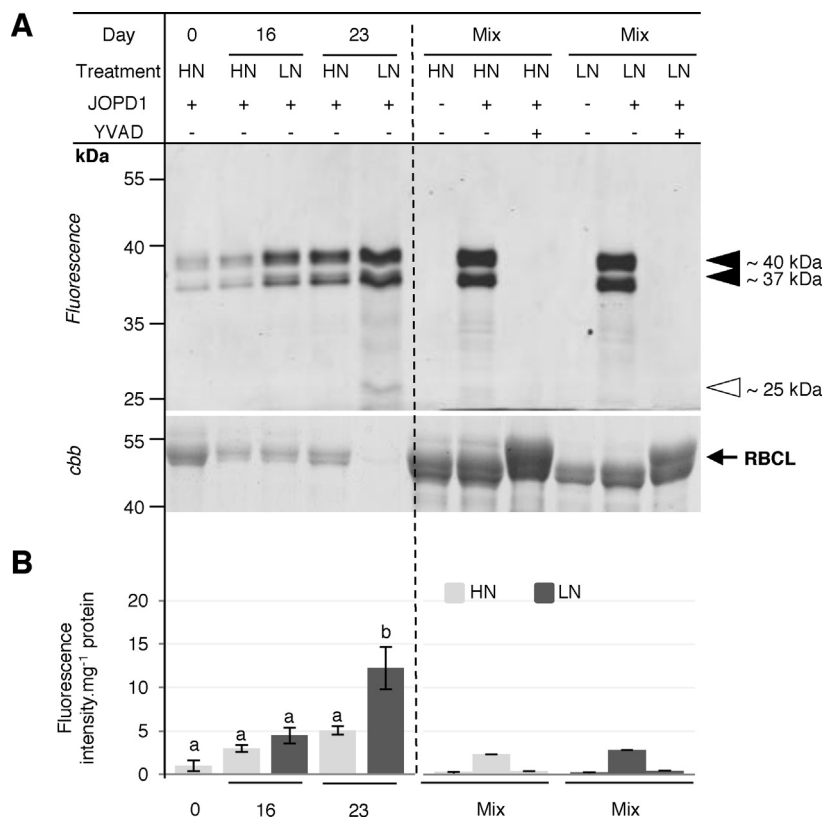


**Fig. 6.** Detection of up-regulated PLCP activities during leaf senescence of *B. napus* L. supplied with low (LN) nitrate for 23 days. Soluble proteins were extracted after 0, 16 and 23 days of HN (3.75 mM  $\text{NO}_3^-$ ) or LN (0.375 mM  $\text{NO}_3^-$ ) treatment and were subjected to standard protease activity profiling with MV201 (specific fluorescent probe of PLCPs) (pH 5.5; 4 h labelling). These extracts were separated by SDS-PAGE and scanned to detect fluorescence (A). The Coomassie brilliant blue-stained (cbb) protein gel shows the total amount of loaded proteins after incubation. Mix corresponds to a mix of 0, 16 and 23 days extracts (HN or LN) in the presence of MV201 alone, MV201 and E64 (specific inhibitor of PLCPs) or in the absence of MV201 (no probe control). The fluorescence intensity representative of the PLCP activity was calculated relative to the protein amount (B). Black arrowheads correspond to proteases already active at day 0 while the white arrowhead shows induced proteases. The gel presented in this figure is representative of three biological replicates. Vertical bars indicate  $\pm$  SE of the mean of three biological replicates. Statistics are represented by letters ( $p < 0.05$ , ANOVA, Newman–Keuls test).

petition experiments consisting of a pre-treatment with 50  $\mu\text{M}$  of E-64, ac-YVAD-cmk, epoxomicin or DFP for 30 min before adding probes MV201, JOPD1, MVB072 or FP-Rh respectively was carried out in parallel aliquots to control the specificity of probes. Proteins were precipitated after labelling by adding 1 mL of ice-cold acetone and centrifuging (15 min, 16,000  $\times g$ , 4 °C). The pellet was dissolved in 2X SDS-PAGE gel-loading buffer, heated at 90 °C for 10 min and separated on 12% SDS-PAGE gels. Labelled proteins were visualised by *in-gel* fluorescence scanning using a Typhoon 9400 scanner (GE Healthcare Life Science) with excitation and emission wavelengths at 532 and 580 nm respectively. Fluorescence signals were quantified by ImageJ software. Then, gels were stained with Coomassie Brilliant Blue stain (0.5 g CBB G250, 10% acetic acid, 45% methanol in ultra-pure water (v/v)), destained (10% acetic acid, 40% methanol in ultra-pure water (v/v)) and scanned to control the protein quantity. Fluorescence signals were used to determine specific activity (expressed as fluorescence intensity  $\text{mg}^{-1}$  protein).

In order to characterize the proteases observed on gels by the fluorescent bands, a pull-down of biotinylated proteins was performed after labelling with a biotin-tagged probe. For this, 700  $\mu\text{g}$  of protein was labelled with 5  $\mu\text{M}$  of DCG04 or FP-biotin in labelling buffer (50 mM NaAc pH 5.5, 2 mM DTT for DCG04 or 50 mM Tris-buffer, pH 7.5 for FP-biotin). Samples were incubated at room temperature in the dark under gentle agitation for 4 h (for DCG04) or 1 h (for FP-biotin). As control, another aliquot of the same sample was treated as described above without probes.

The labelling reaction was stopped and the biotin-proteins were purified using streptavidin beads as described by Chandrasekar et al. [44]. The eluted proteins were separated on 12% SDS-PAGE gels and the protein gels were stained overnight with SYPRO Ruby (Life Technologies). Proteins were detected by scanning the gel at an excitation wavelength of 460 nm in ImageQuant LAS 4000 scanner (Filter Y515 Long Pass 500–520 nm; GE Healthcare Life Sciences). Bands were excised and treated with trypsin as described by Kaschani et al. [45]. Then, tryptic peptides were desalting using Stage Tips C8 (Thermo scientific, Bremen, Germany) according to the manufacturer's instructions. Finally, peptide samples were analysed in triplicate by nano-liquid chromatography tandem mass spectrometry (nano-LC–MS/MS) using a Nano-Acquity-UPLC (C18 column of 75  $\mu\text{m} \times 250$  mm, 1.7  $\mu\text{m}$  particle size; Waters) coupled to an Orbitrap Elite tandem mass spectrometer (Thermo Scientific) with a resolution of 120,000 full-width half maximum at mass/charge 400, Top 20 precursor ion selection, and fragmentation performed in collision-induced dissociation (CID) mode. The samples were loaded in 99.5% buffer A (0.1% formic acid in  $\text{H}_2\text{O}$ ). The gradient used to elute the peptides was started with a 3 min isocratic gradient composed of 3% buffer B (0.1% formic acid in  $\text{CH}_3\text{CN}$ ) followed by a linear gradient from 3–40% of buffer B for 60 min at a flow rate of 250 nL  $\text{min}^{-1}$  and two washes with 97% of buffer B for 3 min. The total length of the analysis was 100 min to allow column re-equilibration.



**Fig. 7.** Activity of VPEs is up-regulated during leaf senescence of *B. napus* L. supplied with low (LN) nitrate for 23 days. Soluble proteins were extracted after 0, 16 and 23 days of HN (3.75 mM NO<sub>3</sub><sup>-</sup>) or LN (0.375 mM NO<sub>3</sub><sup>-</sup>) treatment and were subjected to standard protease activity profiling with JOPD1 (specific fluorescent probe of VPEs) (pH 5.5; 4 h labelling). These extracts were separated by SDS-PAGE and scanned to detect fluorescence (A). The Coomassie brilliant blue-stained (cbb) protein gel shows the total amount of input proteins after incubation. Mix corresponds to a mix of 0, 16 and 23 days extracts (HN or LN) in the presence of JOPD1 alone, JOPD1 and YVAD (inhibitor of VPEs) or in the absence of JOPD1 (no probe control). The fluorescence intensity representative of the VPE activities was calculated relative to the protein amount (B). Black arrowheads correspond to proteases already active at day 0 while the white arrowhead shows induced proteases. The gel presented in this figure is representative of three biological replicates. Vertical bars indicate  $\pm$  SE of the mean of three biological replicates. Statistics are represented by letters ( $p < 0.05$ , ANOVA, Newman–Keuls test).

Raw MS data were converted into mgf files using MS convert and processed using Mascot software and the following parameters: the error tolerance was fixed at 10 ppm for the precursor ion and at 0.5 Da for the fragment ion. The enzyme used was trypsin and only one missed cleavage was allowed. The post-translational modifications search setting was as follows: fixed modification for cysteine carbamidomethyl and variable modification for asparagine and glutamine deamidation and methionine oxidation. The protein search was performed using the *B. napus* 20150610 database (101,040 sequences; 33,618,840 residues). The false discovery rate was estimated empirically from decoy hits; identified proteins were filtered to an estimated 1% FDR (false discovery rate). The assigned protein of best match is provided alongside the GenBank accession number. Score, peptide matches, significant peptide matches (Match(sig)), experimental mass and theoretical mass are also presented. Protein sequences were also matched against sequences of *Brassicaceae* proteins using the NCBI Blast Protein Database (algorithm blastp) and better results of blasts are presented with the name of the protein, the organism, the gene accession number and the percentage of sequence identity. Finally, PLCPs were classified according to the classification of Richau et al. [37] while SPs were classified according to the MEROPS database.

## 2.5. Statistical analysis

For all measurements, three biological repeats were analysed ( $n=3$ ). All the data are presented as the mean  $\pm$  standard error (SE). The normality of the data was studied with the Ryan–Joiner test at 95%. Analysis of variance (ANOVA) and the Newman–Keuls

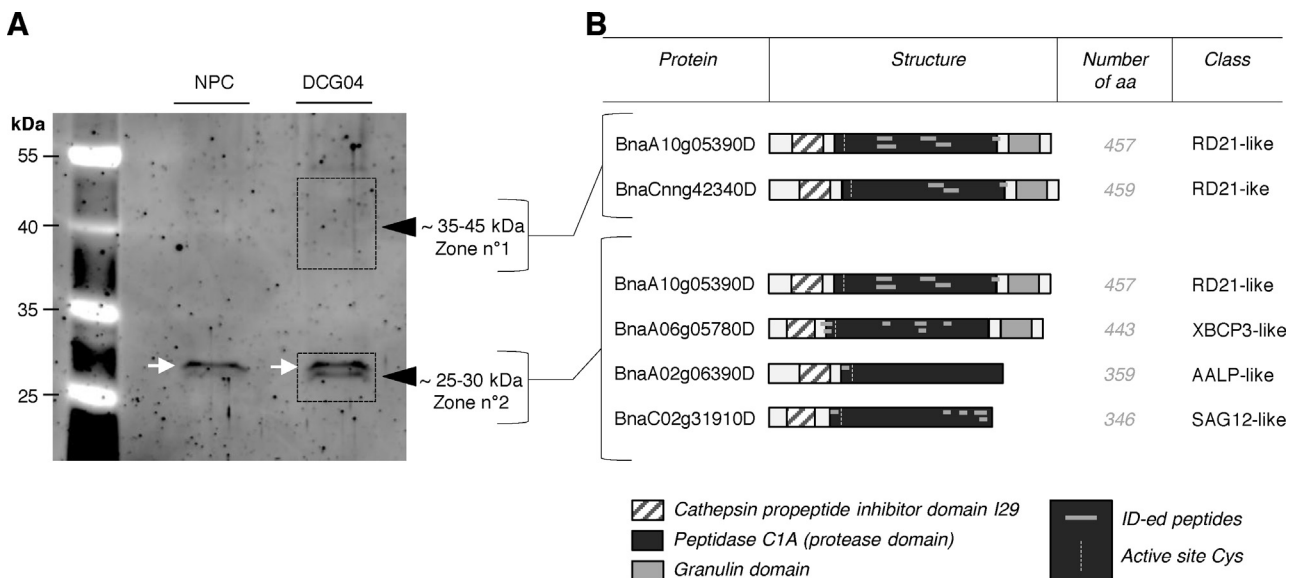
test were used to compare the means by using Microsoft® Excel 2010/XLStat® 2014. When the normality law of the data was not respected, the non-parametric test of Kruskal–Wallis was carried out. Statistical significance was postulated at  $p < 0.05$ .

## 3. Results and discussion

In order to characterize key mechanisms that could assist in improving the protein recycling and the leaf NRE of oilseed rape, the objective was to identify senescence-associated proteases (SAPs) implicated in protein degradation during leaf senescence of genotype Aviso, which was previously characterized for its high leaf NRE in response of nitrate limitation [17].

### 3.1. Leaf senescence and soluble protein degradation

To study the changes in physiological and biochemical parameters associated with senescence in a mature leaf of oilseed rape, six week old plants were subjected to ample (HN: 3.75 mM NO<sub>3</sub><sup>-</sup>) or low nitrogen supply (LN: 0.375 mM NO<sub>3</sub><sup>-</sup>) for 23 days. Mature leaves at rank no. 12 (L12) were chosen for the study of senescence progression via the analysis of chlorophyll content, quantity of soluble proteins and protease activities. As expected, a limitation of nitrate (LN) accelerated the leaf senescence processes compared to plants subjected to an ample nitrate supply (HN). Indeed, chlorophyll content was significantly lower after 23 days under LN conditions (22.3  $\pm$  4 SPAD units) compared to HN supply (39.8  $\pm$  3) (Fig. 1A). Because leaf redness through anthocyanin accumulation is commonly considered as a symptom of abiotic stresses



**Fig. 8.** Identification of PLCPs labelled with DCG04 in senescent leaf of *B. napus* L. after 23 days of LN treatment. To characterize the cysteine proteases observed by fluorescent bands on gels (Fig. 6), a pull-down of biotinylated proteins was prepared after labelling with the biotin-tagged probe DCG04 (A). The experiment was performed only on soluble protein extracts from L12 of plants subjected to LN supply after 23 days. The labelling reaction was stopped and the biotin-proteins were purified using streptavidin beads. The eluted proteins were separated on 12% SDS-PAGE gel and stained with SYPRO Ruby. Proteins were detected by scanning the gel at an excitation wavelength of 460 nm in an ImageQuant LAS 4000 scanner. NPC corresponds to the no probe control. Streptavidin beads are indicated by white arrows. Black arrowheads correspond to zones no. 1 (35–45 kDa) and no. 2 (25–30 kDa), which were excised and analysed by LC-MS/MS. (B) Identified proteases for each zone and their structures. All protease domains are preceded by a cathepsin propeptide inhibitor domain (I29). RD21-like and XBCP3-like proteases also carry a C-terminal extension, containing a granulin domain. Tryptic fragments identified (ID-ed peptides) are indicated in the proteins with gray bars and white bars corresponding to the active site cysteine. Finally, proteins were classified according to the classification of Richau et al. [37].

especially in case of mineral limitation such as sulfur, phosphorus and N deficiency in many plants [46–50] including oilseed rape [41], we have studied the changes of anthocyanins in the mature leaf of plants subjected to ample or limited nitrate supply. The level of anthocyanins in L12 had tripled during the 23 days of LN treatment but only doubled under HN treatment (Fig. 1B). These results confirm that N limitation can cause a rapid induction of anthocyanins in senescing leaves of oilseed rape. Acting as a “sunscreen” in case of excess light and as scavengers for reactive oxidative species, an accumulation of anthocyanins in leaves may protect the photosynthetic apparatus from photodamage and facilitate the recovery of nutrients, especially N, from senescing leaves of oilseed rape as previously reported in other plants [51,52,46]. Furthermore, after 23 days, the nitrate limitation induced the strong degradation of soluble proteins in L12 (~76%) versus only ~35% for L12 of HN plants (Fig. 1C), as described previously for the genotype Aviso [17].

As the N need of growing organs is mainly fulfilled by the degradation of RuBisCO during leaf senescence [20,21], we studied, in particular, the degradation of RBCL (Large subunit of RuBisCO) *in vitro* at pH 5.5 or 7.5. At pH 5.5 (Fig. 2), the degradation of RBCL by proteases present in extracts of L12 increased at D16 and D23 compared to D0 regardless of the treatment. Nevertheless, after 23 days of nitrate limitation, the RBCL was more degraded ( $-89.4 \pm 3.4\%$ ) than under HN supply ( $-46.9 \pm 9\%$ ). Otherwise, no degradation of RBCL was observed at pH 7.5 regardless of the date or treatment (data not shown). These results suggest that RBCL degradation in oilseed rape might occur outside the plastid in cellular compartments with an acidic pH such as SAVs (Senescence Associated Vesicles) or in the central vacuole through the transfer by RCBs (RuBisCO containing bodies) [53–57]. Other studies have shown that chloroplast proteins such as RuBisCO can be partially degraded in the chloroplast itself by chloroplastic proteases [19,58,59]. Even if chloroplasts are unable to carry out complete breakdown of the RBCL, it has been reported that a 44 kDa fragment from cleaved RBCL accumulates in chloroplasts isolated from

senescing wheat leaves [55,58]. Other chloroplastic proteins such as LHCII can be completely degraded in the plastid at pH 7.5 [55,60], and novel chloroplast vesiculation-containing vesicles (CCVs) have been identified and implicated in chloroplast degradation in *Arabidopsis* [61]. CCVs containing chloroplast proteins were released from chloroplasts and mobilized to the vacuole for degradation through a pathway that is independent of autophagy and SAVs.

### 3.2. Senescence-associated protease (SAP) activities in oilseed rape leaves

To determine which protease classes could be involved in the degradation of soluble proteins during senescence (Figs. 1C and 2), we studied protease activities *in vitro* at acidic pH (5.5) or neutral pH (7.5) during leaf senescence of plants subjected to N limitation (LN) compared to plants with ample nitrate supply by two different methods. The first method consisted in studying the RBCL degradation only at acidic pH (analysis of RBCL degradation was not performed at neutral pH because of the low contribution of proteolytic activities at this pH, see Section 3.1) in the presence or absence of known inhibitors of specific protease classes. Then, in order to identify active proteases at pH 5.5 and pH 7.5, we performed in parallel a protease activity profiling during leaf senescence using activity-based probes specific for different protease classes [36]. Results indicated that many classes of proteases were active during leaf senescence of oilseed rape, including the proteasome, aspartic, serine and cysteine proteases.

#### 3.2.1. Aspartic proteases (APs)

At pH 5.5, the presence of pepstatin A (inhibitor of aspartic proteases) at D0 inhibited RBCL degradation by  $45.2 \pm 6\%$  at D0 but this was not increased at D16 or D23 under the HN or LN treatments (Fig. 3A). These results showed that APs seem to be active at pH 5.5 during senescence but these activities are not affected by an acceleration of senescence in LN plants. This is in agreement

**Table 3**LC-MS/MS identification of cysteine proteases labelled with the probe DCG04 in senescent leaf of *B. napus* L. after 23 days of nitrate limitation.

Cutting zone (kDa)	Protein accession no. [Brassica napus]/NCBI accession no.	Score	Matches	Match (sig)	Exp. Mass (Da)	Theo. Mass (Da)	Result of BLAST protein [Brassica napus]-protein [organism]/NCBI accession no./% identity	Classification of Richau et al. [37]
35–45	BnaA10g05390D [Brassica napus]/gi 674926680	201	14	9	51650	50484.76	Senescence-associated cysteine protease [Brassica oleracea]/gi 18141285/ 92% Cysteine proteinase RD21a [Arabidopsis thaliana]/gi 18401614/ 83%	RD21A-like
	BnaCnng42340D [Brassica napus]/gi 674871100	65	5	4	51828	50662.93	Senescence-associated cysteine protease [Brassica oleracea]/gi 18141285/ 86% Cysteine proteinase RD21a [Arabidopsis thaliana]/gi 18401614/ 82%	RD21A-like
25–30	BnaA10g05390D [Brassica napus]/gi 674926680	60	5	4	51650	50484.76	Senescence-associated cysteine protease [Brassica oleracea]/gi 18141285/ 92% Cysteine proteinase RD21a [Arabidopsis thaliana]/gi 18401614/ 83%	RD21A-like
	BnaA06g05780D [Brassica napus]/gi 674939790	52	8	4	50101	48821.05	PREDICTED: low-temperature-induced cysteine proteinase [Brassica rapa]/gi 685315355/ 99% Papain-like cysteine peptidase XBCP3 [Arabidopsis thaliana]/gi 614600257/89%	XBCP3-like
	BnaA02g06390D [Brassica napus]/gi 674924155	40	2	1	39682	39193.26	PREDICTED: thiol protease aleurain-like [Brassica rapa]/gi 685271156/ 96% Senescence-associated cysteine protease [Brassica oleracea]/gi 18141289/ 90% Thiol protease aleurain [Arabidopsis thaliana]/gi 18424347/ 89%	AALP-like
	BnaC02g31910D [Brassica napus]/gi 674945739	38	7	1	38506	38079.10	Cysteine protease SAG12 [Arabidopsis thaliana]/gi 18422605/ 86%	SAG12-like

To characterize the cysteine proteases observed by fluorescent bands on gels (Fig. 6), a pull-down of biotinylated proteins was prepared after labelling with biotin-tagged probe DCG04 and detected zones were excised, treated and analysed by LC-MS/MS (for details see Section 2). LC-MS/MS spectra were searched against the *Brassica napus* L. database using MASCOT 2.5.0 and only scores higher or equal to 30 were considered. The assigned protein of best match is provided with the GenBank accession number. Score, peptide matches, significant peptide matches (Match(sig)), experimental mass and theoretical mass are also presented. Protein sequences were also matched against sequences of *Brassicaceae* proteins using the NCBI Blast Protein Database (algorithm blastp) and better results of blasts were presented with the name of the protein, the organism, the gene accession number and the percentage of sequence identity. Finally, proteins were classified according to the classification of Richau et al. [37].

with previous proteomic studies reporting that the abundance of an aspartic protease was maintained to a high level during leaf senescence of *B. napus* [25]. Several studies have demonstrated that the aspartic protease CND41 may play a crucial role during leaf senescence [14,26,27]. For instance, in tobacco leaves, CND41 has a high activity at acidic pH and this protease was able to degrade RBCL *in vitro* [26]. The present study supports a role for APs during leaf senescence in oilseed rape, but this might be underestimated because pepstatin A does not inhibit tobacco CND41 [26].

### 3.2.2. The proteasome

Because the proteasome is physiologically active under neutral pH, the proteasome activity was studied at pH 7.5. In our experiment, RBCL was not degraded at pH 7.5 (data not shown) indicating that, at neutral pH, the proteasome is not implicated in the degradation of RBCL. To detect the active proteasome during senescence, an activity-dependent labelling using MVB072 (a specific probe of the proteasome) was performed (Fig. 4, [40]). Labelling with MVB072 allowed the detection of the three catalytic subunits of proteasome:  $\beta 1$ ,  $\beta 2$  and  $\beta 5$  (Fig. 4A). Quantification of the sum of activities of these subunits demonstrated that global proteasome activity remained stable during leaf senescence in oilseed rape supplied with ample or low nitrate (Fig. 4B). A similar result was found in dark-induced detached senescent leaves of wheat, which have a constant activity of the 20S proteasome [62]. Several studies also suggest that the proteasome remains functional until late stages of senescence [23]. However, a study of proteomic events associated with N remobilization during leaf senescence of oilseed rape demonstrated that the  $\beta 1$  subunit was accumulated upon N limitation or starvation [25]. These data suggest that the proteasome plays an important role during senescence, not particularly in the remobilization of Rubisco, but for example to limit the production of reactive oxygen species and/or to contribute to the degradation of other damaged proteins.

### 3.2.3. Serine proteases (SPs)

Because SPs are located in the vacuole [63] and in the chloroplast [23,64], SP activities were studied using (i) RBCL as substrate, under acidic pH with or without the SP inhibitor aprotinin (Fig. 3B) and (ii) a standard protease activity profiling of SPs using FP-Rh, a specific fluorescent probe of serine proteases performed at pH 7.5 (Fig. 5).

The presence of aprotinin (inhibitor of SPs) at D0 inhibited RBCL degradation by 22.4%, but this was not increased at D16 or D23 under the HN or LN treatments (Fig. 3B). It seemed that SPs were physiologically active under acidic pH and were able to degrade RBCL but their contribution was relatively low when compared to other protease classes (Fig. 3). A study on leaves of naturally senescent wheat also suggested that SPs participate in RBC degradation and N remobilization [65]. Otherwise, our data suggest that at neutral pH, SPs do not contribute significantly to RBCL proteolysis during leaf senescence of oilseed rape, irrespective of the nitrate supply.

To verify if SPs were active at neutral pH, an activity-dependent labelling using FP-Rh (a specific probe of SPs) was carried out at pH 7.5 (Fig. 5). Many labelled serine hydrolases were detected after labelling with FP-Rh at pH 7.5 (Fig. 5A). Fluorescent bands were detected at ~70, ~40–50, ~38, ~35 and ~25–30 kDa and these signals were absent in the no-probe controls and suppressed upon pre-treatment with DFP. Quantification demonstrated that global labelling of serine hydrolases was up-regulated during leaf senescence in response to a limitation of nitrate (Fig. 5B). This was due to an increase in serine hydrolase labelling between D0 and D23 at ~38–50 kDa ( $p = 0.007$ ), ~35 kDa ( $p = 0.021$ ) and ~25–30 kDa ( $p = 0.035$ ) (data not shown). In response to nitrate limitation, labelling of serine hydrolases at ~70 kDa also increased between D0 and D16.

To identify the labelled SPs, a pull-down of biotinylated proteins from L12 of plants subjected to LN supply over 23 days was carried out after activity-dependent labelling using the biotin-tagged FP probe. Several labelled SPs were identified from signals at 70 and 35–40 kDa (Fig. S2 and Table 2). SPs at 70 kDa were identified as four different subtilisins (S8) and seven prolyl oligopeptidase-like proteases (POPLs, S9) (Table 2). In *A. thaliana*, activity-dependent labelling with similar probes also permitted the identification of POPLs (S9) and subtilase-like (S8) proteases at 70 kDa [36,66,67,68]. Some studies have indicated a role for subtilases during natural senescence in wheat and barley leaves [24,63,65]. In addition, during the programmed cell death (PCD) induced by vicerin or heat shock treatment, subtilisins (particularly active at neutral pH) are required for RuBisCO cleavage [69]. These results may indicate that the RBCL is not degraded at pH 7.5 *in vitro*. Thus, RBCL does not seem to be the direct target of SPs that are active at neutral pH, in contrast to those active at acidic pH. Interestingly, BnaA03g44620D corresponds to tripeptidyl peptidase II (TPPII), which degrades peptides released by the proteasome [70]. In parallel, four carboxylesterases (CXEs) were identified at both 70 and 35–40 kDa while the three methylesterases (MESs) in *B. napus* correspond to the methylesterase 9 of *A. thaliana* (AtMES-9). MESs can hydrolyze methylsalicylate, methyljasmonate, and methylindoleacetic acid and may play a role in the regulation of these plant hormones [71,72]. Additional experiments are required to characterize the targets of these SPs to determine their role in the leaf senescence of oilseed rape.

### 3.2.4. Cysteine proteases (CPs): PLCPs and VPEs

CP activities were studied under acidic pH (5.5) because these proteases are located in the vacuole or in SAVs [30,54]. CP activities were studied firstly using RBCL as substrate, with or without E-64 (inhibitor of papain like cysteine proteases (PLCPs)) or Ac-YVAD-cmk (inhibitor of vacuolar processing enzymes (VPEs)) (Figs. 3C and D). Additionally, a standard protease activity profiling of PLCPs and VPEs using the probes MV201 or JOPD1, specific fluorescent probes of PLCPs and VPEs respectively, were performed at pH 5.5 (Figs. 6 and 7).

The inhibition of RBCL degradation due to E-64 (inhibitor of PLCPs) was significantly more effective at D23 (78.8% inhibition (HN); 79% (LN)) than at D0 (11.9%) (Fig. 3C). The degradation of RBCL was also significantly inhibited by Ac-YVAD-cmk (inhibitor of VPEs) after 16 or 23 days in L12 of HN and LN plants (Fig. 3D). These results show that the increase in RBCL degradation during senescence (Fig. 2) correlates with a significant increase in PLCP and VPE activities at pH 5.5 focused on the RBCL degradation (Figs. 3C and D). Otherwise, this increase was not significantly different between HN and LN samples.

The results obtained by protease activity profiling at pH 5.5 with MV201 (specific fluorescent probe for PLCPs) or JOPD1 (specific fluorescent probe for VPEs) are presented in Figs. 6 and 7, respectively. Many PLCPs were detected after labelling with MV201 at pH 5.5 (Fig. 6A): at ~40, ~30–35 and ~27 kDa and these signals were absent in the no-probe controls and suppressed upon pre-incubation with PLCP inhibitor E-64. In response to nitrate limitation, the total PLCP labelling was significantly up-regulated in L12 of LN plants after 23 days of treatment (Fig. 6B). Interestingly, one additional band at ~27 kDa appeared only after 23 days of nitrate limitation. Labelling with JOPD1 at pH 5.5 displayed active VPEs at ~40, ~37 and ~25 kDa and these signals were absent in the no-probe controls and suppressed upon pre-incubation with the VPE inhibitor, Ac-YVAD-cmk (Fig. 7A). VPE activities were significantly up-regulated in L12 of plants submitted to LN treatment after 23 days (Fig. 7B). Moreover, one additional fluorescent signal appeared at ~25 kDa after 23 days in response to LN treatment (Fig. 7A). These results reveal that the increased degradation of soluble proteins

such as RBCL under LN correlates with an increase in several PLCP and VPE activities at pH 5.5 (Figs. 6 and 7).

It has already been shown that CPs are involved in the degradation of stromal proteins, particularly RBC, during senescence in wheat and tobacco leaves [21,31,55]. A study of dark-induced senescent tobacco leaves showed that RBC degradation in isolated SAVs was blocked by E-64 [57]. More specifically, some PLCPs were up-regulated during senescence in sweet potato, soybean and barley leaves [63,73–75]. In addition, transcript and protein levels of PLCP SAG12 are up-regulated during leaf senescence of oilseed rape subjected to a limitation of N [25]. These data strongly indicate that PLCPs can be involved in proteolysis associated with leaf senescence in oilseed rape.

In order to identify the labelled PLCPs, a pull-down of biotinylated proteins from L12 of plants subjected to LN supply for 23 days was performed after activity-dependent labelling using the biotin-tagged DCG04 probe and allowed us to identify five PLCPs (Fig. 8 and Table 3) corresponding to the fluorescent band observed at 27 kDa using MV201 labelling (Fig. 6A).

Two of these PLCPs were classified as RD21A-like proteases according to the classification of Richau et al. [37] and were identified as close homologues of RD21A from *A. thaliana* in the active form at 25 kDa (BnaA10g05390D) and in the intermediate active isoform at 40 kDa (BnaA10g05390D and BnaCnng42340D). RD21 is a PLCP expressed in leaves during senescence and its activity increase in senescent leaves of *A. thaliana* [76,77]. These results strongly suggested that several isoforms of *B. napus* proteins may correspond to RD21A in *A. thaliana*. This protease is synthesized as an inactive pro-protease that is cleaved to the mature/active form in the vacuolar acidic environment, and it has been proposed that the pro-protease might be contained inside ER-bodies with a non-optimum pH that keeps the protease inactive [76,78].

The three other PLCPs were identified as XBCP3-like, AALP-like and SAG12-like proteases according to the classification of Richau et al. [37]. Interestingly, BnaA02g06390D (AALP-like protease) is a close homologue of the senescence-associated cysteine protease, BoCP5, an aleurain-like protein in *Brassica oleracea* (gi|18141289). In broccoli florets the suppression of BoCP5 delays floret senescence [79]. Consequently, we strongly suspect that BnaA02g06390D in oilseed rape has the same proteolysis function during leaf senescence as BoCP5 in *B. oleracea*. Finally, BnaC02g31910D is homologous to SAG12 of *A. thaliana*. SAG12 is a senescence-associated marker gene [80] and encodes for the protease SAG12, which was found specifically in the SAVs [30]. Moreover, Desclos et al. [25] showed that in *B. napus* leaves subjected to a nitrate limitation, SAG12 expression and SAG12 protein abundance are up-regulated during senescence. Consequently, these results indicate that PLCPs related to RD21A or SAG12 proteases could play a crucial role in the efficient proteolysis associated with leaf senescence of *B. napus*, especially in response to N limitation.

In parallel, VPEs are not implicated in the degradation of chloroplastic proteins such as RBC, but they are responsible for the maturation of several vacuolar proteins [81]. In *Arabidopsis* leaves, it was suggested that  $\alpha$ VPEs and  $\gamma$ VPEs (specific to the vegetative organs) regulate the activation of proteins in the lytic vacuole [82]. For instance, it was reported that SAG2 and RD21 genes were induced during leaf senescence and correlated with the induction of VPEs [82]. Previous work has also suggested that  $\gamma$ VPE might activate proteases involved in amino acid recycling during the senescence of *A. thaliana* leaves [33]. Because VPEs might be responsible for the maturation/activation of other proteases such as PLCPs, the increase in VPE activities observed in our study (Figs. 3D and 7) could contribute to an increase in PLCP activities (Figs. 3C and 6). However, some PLCPs, such as RD21A in *Arabidopsis* [83],

do not require VPEs for their activation, so their role in senescence remains unclear.

#### 4. Concluding remarks

In conclusion, many classes of proteases are implicated in protein degradation during natural senescence or senescence induced by nitrate limitation in *B. napus* leaves at the vegetative stage. Aspartic proteases, serine proteases, the proteasome and cysteine proteases are highly active at different pH values associated with different cellular compartments. A limitation of nitrate availability, characterized by an acceleration of senescence processes, leads to an increase in proteolytic activities caused by serine proteases and particularly cysteine proteases (both PLCPs and VPEs). As soluble protein degradation in senescent leaves of oilseed rape is crucial for the improvement of N remobilization efficiency (NRE), characterization of protease activities is a key for the comprehension of leaf N remobilization. Moreover, the genotypic differences of leaf NRE observed in response to nitrate limitation [17] might be associated with different or contrasted protease activities during leaf senescence. This study also revealed that activity-based protein profiling is a relevant method for future investigations of genotypic variability of leaf proteolysis efficiency and characterization of protease machinery associated with a high leaf NRE in winter oilseed rape.

#### Author contributions

Marine Poret and Jean-Christophe Avice contributed to the experimental design and tissue sampling. Marine Poret and Balakumaran Chandrasekar carried out the protease activity profiling using activity-dependent fluorescent probes. Marine Poret performed other biochemical measurements and proteases analyses, statistical analyses, interpretation of data and drafting the article. Marine Poret, Renier van der Hoorn and Jean-Christophe Avice were involved in revising the manuscript for important intellectual content.

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#### Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.plantsci.2016.02.011>.

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