Italian legumes: effect of sourdough fermentation on lunasin-like polypeptides

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Abstract

Background: There is an increasing interest toward the use of legumes in food industry, mainly due to the quality of their protein fraction. Many legumes are cultivated and consumed around the world, but few data is available regarding the chemical or technological characteristics, and especially on their suitability to be fermented. Nevertheless, sourdough fermentation with selected lactic acid bacteria has been recognized as the most efficient tool to improve some nutritional and functional properties. This study investigated the presence of lunasin-like polypeptides in nineteen traditional Italian legumes, exploiting the potential of the fermentation with selected lactic acid bacteria to increase the native concentration. An integrated approach based on chemical, immunological and ex vivo (human adenocarcinoma Caco-2 cell cultures) analyses was used to show the physiological potential of the lunasin-like polypeptides.

Results: Italian legume varieties, belonging to Phaseolus vulgaris, Cicer arietinum, Lathyrus sativus, Lens culinaris and Pisum sativum species, were milled and flours were chemically characterized and subjected to sourdough fermentation with selected Lactobacillus plantarum C48 and Lactobacillus brevis AM7, expressing different peptidase activities. Extracts from legume doughs (unfermented) and sourdoughs were subjected to western blot analysis, using an anti-lunasin primary antibody. Despite the absence of lunasin, different immunoreactive polypeptide bands were found. The number and the intensity of lunasin-like polypeptides increased during sourdough fermentation, as the consequence of the proteolysis of the native proteins carried out by the selected lactic acid bacteria. A marked inhibitory effect on the proliferation of human adenocarcinoma Caco-2 cells was observed using extracts from legume sourdoughs. In particular, sourdoughs from Fagiolo di Lamon, Cece dell’Alta Valle di Misa, and Pisello riccio di Sannicola flours were the most active, showing a decrease of Caco-2 cells viability up to 70%. The over-expression of Caco-2 filaggrin and involucrin genes was also induced. Nine lunasin-like polypeptides, having similarity to lunasin, were identified.

Conclusions: The features of the sourdough fermented legume flours suggested the use for the manufacture of novel functional foods and/or pharmaceuticals preparations.

Keywords: Legumes, Sourdough, Lactic acid bacteria, Lunasin
of pulses is recently associated with protective or therapeu-
tic effects on chronic health conditions, such as cardio-
vascular diseases, diabetes, cancer, overweight, and obesity [2]. Owing to the low cost and easy adaptation to
grow under poor conditions, pulses are used as staple
foods in several low-income countries, serving as main
source of both protein and calories [2, 3]. Otherwise, in
high-income countries of America and Europe, pulses
consumption is low and efforts are done to promote their
healthy intake.

Several health organizations recommend pulse con-
sumption as a part of a healthy diet and initiatives are
addressed to increase the cultivation, intake, and food
processing uses [2, 4]. The protein content of pulses
ranges from 20 to 40 % of dry weight. Within this inter-
val, the most abundant are the seed storage proteins.
The remaining part are minor or housekeeping proteins,
which include enzymes, protease, amylase inhibitors, lec-
tins, lipoygenases, defense proteins, and others [3, 5].
Legume seeds also contain proteins, which are consid-
ered as anti-nutritional compounds due to the effect on
the quality of the diet. Nevertheless, the harmful effects
of such compounds is easily inactivated after cooking
or processes like fermentation, germination and dehull-
ing [6]. Once inactivated, lectins or protease inhibitors
may present potential health benefits. Protease inhibitors
are potential anti-inflammatory and anticancer agents,
whereas lectins have demonstrated to play a key role in
preventing certain cancers and activating innate defense
mechanisms. Besides, lectins are also proposed as thera-
pic agents to prevent or control obesity [5].

Many of the physiological and functional properties
of proteins are attributed to biologically active peptides
which are often encrypted in the native sequence [7, 8].
Biogenic or bioactive peptides are released from their
precursor proteins either by enzymes during gastroin-
testinal digestion or through proteolysis (e.g., microbial
fermentation), which occurs during food processing [8].
In particular, legume hydrolysates and bioactive peptides
had in vitro activities towards cancer and cardiovascular
diseases or their physiological manifestations like oxida-
tive damage, inflammation, hypertension, and high
cholesterol [2]. Lunasin is a 43-amino acid peptide with
anticancer, anti-inflammatory, antioxidant and choles-
terol lowering activities [9]. It is purified from soybean
and commercialized as an ingredient or dietary supple-
ment. In the quest for readily available natural sources
of lunasin, the identification and purification of lunasin
from different vegetable sources deserve a marked inter-
est [10, 11]. Moreover, the potential of sourdough fer-
m entation for increasing the concentration of lunasin
in food matrices was recently investigated [12]. Sour-
dough is the natural starter traditionally used for making
leavened baked goods, harboring a rich lactic acid bacte-
ria and yeast microbiota. A large number of studies [13]
showed that the fermentative and proteolytic activities
of sourdough lactic acid bacteria not only determined opti-
mal sensory, technology and nutritional characteristics,
but also increased the functional value of leavened baked
goods. Compared to unfermented soybean, amaranth,
barley and wheat flours, the concentration of lunasin and
related fragments increased up to four times during fer-
m entation with lactic acid bacteria, which were selected
based on proteolytic activities [12].

This study reported the presence of lunasin-like poly-
peptides in nineteen traditional Italian legumes, and
exploited the potential of the fermentation with selected
lactic acid bacteria to increase the native concentration.
An integrated approach based on chemical, immuno-
logical and ex vivo (human adenocarcinoma Caco-2 cell
cultures) analyses was used to show the physiological
potential of the lunasin-like polypeptides.

Results
Chemical and microbiological characteristics of the flours
All the traditional Italian legumes used in this study
have specific certifications (names, abbreviations, geo-
 graphical origin and certification are listed in Table 1).
Legume grains were milled to obtain the correspond-
ing flours. The proximate composition of the flours is
reported in Table 2. Moisture ranged from 7.2 ± 0.4 to
11.1 ± 0.7 %. The protein concentration of all the flours
was higher than 15 % of dry matter (d.m.). In particular,
grass pea flours (CS and CC) showed the highest values
(>24.0 %). The lipid concentration varied from 1.4 ± 0.2
to 3.5 ± 0.1 % of d.m., with the exception of chickpea
flours (CM and CV), showing a significantly (P < 0.05)
higher concentration. All flours had an amount of car-
bohyd r ate s higher than 60 % of d.m. The highest concen-
tration was found for FCo and LA flours, produced by
milling Fagiol o di Controne kidney beans and Lentichia
di Altamura lentils, respectively. Legume flours had con-
centrations of total dietary fiber higher than 17 % of d.m.
PS and CC flours had the highest values (35.5 ± 2.9 and
32.1 ± 2.0 %, respectively), whereas the lentils group con-
tained the lowest value of total dietary fiber. Ash ranged
from 2.1 ± 0.2 to 4.6 ± 0.3 %, being the lowest values for
lentil flours.

Total mesophilic aerobic bacteria ranged from
1.61 ± 0.32 to 4.31 ± 0.11 log cfu/g (Table 3). Entero-
bacteria were below 2.0 log cfu/g in all the samples.
The number of presumptive lactic acid bacteria varied
from 1.0 ± 0.08 to 2.54 ± 0.12 log cfu/g. Except for
glass pea flour CC (4.23 ± 0.22 log cfu/g), yeasts were
1.11 ± 0.05–2.77 ± 0.20 log cfu/g. Yeasts were not found
in 10 g of LP, FV, FSc, and FBl flours. Molds were found
at 2.0 ± 0.21–3.51 ± 0.13 log cfu/g in most of the flours and were absent in 10 g of LS, FCo, CC, and PS flours.

Lactic acid bacteria fermentation
After incubation for 24 h at 30 °C, control doughs (D), without bacterial inoculum, showed values of pH that ranged from 5.8 ± 0.2 to 6.5 ± 0.3, corresponding to values of TTA of 3.8 ± 0.2 to 8.5 ± 0.5 ml 0.1 M NaOH/10 g of dough (Table 4). When L. plantarum C48 and L. brevis AM7 were used as starters for sourdough (S) fermentation, the cell density of presumptive lactic acid bacteria was 9.8–10.2 log cfu/g. All S had values of pH significantly (P < 0.05) lower (3.9 ± 0.1–4.5 ± 0.3) than the corresponding D. TTA ranged from 20.2 ± 1.3 to 27.2 ± 1.5 ml NaOH/10 g of dough.

The concentration of total free amino acids (FAA) of D and S is reported in Table 4. Before incubation, doughs had concentrations of total FAA varying from 2429 ± 20 (D made with LU flour) to 4547 ± 18 mg/kg (D made with FBw flour). In several cases, the concentration of total free amino acids (FAA) of S was significantly (P < 0.05) higher than that of the corresponding D. The average increase was ca. 28 % for kidney bean and pea S. Almost the same average increases (23–26 %) were found for grass pea, chickpea, and lentil S. Compared to D, slight increases of total FAA were shown by FSa, CM, CC, LS, and PS sourdoughs.

Western blot
The water/salt soluble extracts (WSE) of D and S were analyzed by SDS PAGE, and the proteins were electroblotted and detected after incubation with lunasin polyclonal primary antibody (Fig. 1). A polyclonal primary antibody was already used for identification and quantification of lunasin and its use proposed for optimized methods of detection [14]. Lunasin peptide was not found in any of the samples analyzed. Nevertheless, different immunoreactive bands, having molecular masses higher than lunasin, were found. Among D, grass pea and chickpea doughs did not show any immunoreactive bands, while pea and all the lentil doughs (with the exception of LV) showed very weak signals, mainly distributed below 15 kDa. A large variability was found among the doughs made with bean flours. Very weak bands were found for Fl, FCo, FCu, and Fst, while a large protein band (molecular mass of ca. 17 kDa) was found for LV, FV, FSa, FBw, and FBb.

Table 1 List and abbreviations of the Italian legumes

| Legume                  | Name                        | Abbreviation | Product certification* | Origin       |
|-------------------------|-----------------------------|--------------|------------------------|--------------|
| Kidney bean (Phaseolus vulgaris) | Fagiolo di Lamon           | FL           | IGP                    | Veneto       |
|                         | Fagiolo di Controne         | FCo          | DOP                    | Campania     |
|                         | Fagiolo di Cuneo            | FCu          | PAT                    | Piedmont     |
|                         | Fagiolo Stregoni            | FSt          | PAT                    | Piedmont     |
|                         | Fagiolo Vellutina           | FV           | PAT                    | Sicily       |
|                         | Fagiolo di Saluggia         | FSa          | PAT                    | Piedmont     |
|                         | Fagiolo Badda di Polizzi (white) | FBw    | SFP                    | Sicily       |
|                         | Fagiolo Badda di Polizzi (black) | FBB     | SFP                    | Sicily       |
| Chickpea (Cicer arietinum)  | Cece di Merella            | CM           | PAT                    | Piedmont     |
|                         | Cece dell’Alta Valle del Misa | CV       | SFP                    | Marche       |
| Grass pea (Lathyrus sativus)  | Cicicheria di Serra de Conti | CS           | SFP                    | Marche       |
|                         | Cicicheria di Campodimele   | CC           | PAT                    | Lazio        |
| Lentil (Lens culinaris)   | Lenticchia di Castelluccio di Norcia | LN     | IGP                    | Umbria       |
|                         | Lenticchia di Ustica        | LU           | SFP                    | Sicily       |
|                         | Lenticchia di Santo Stefano di Sessanio | LS | SFP                    | Abruzzo      |
|                         | Lenticchia rossa di Pantelleria | LP    | PAT                    | Sicily       |
|                         | Lenticchia di Altamura      | LA           | PAT                    | Apulia       |
|                         | Lenticchia di Villalba      | LV           | PAT                    | Sicily       |
| Pea (Pisum sativum)       | Pisello ricco di Sannicola  | PS           | PAT                    | Apulia       |

Product certifications and origin are also reported

* IGP (Indicazione Geografica Protetta, Protected Geographical Indication) and DOP (Denominazione d’Origine Protetta, Designation of Protected Origin) are regulated by Reg. (CE) N. 510/2006 (20.03.2006); PAT (Prodotti Agroalimentari Tradizionali, Traditional Food Products) are included in the list of the Italian Ministry of Agriculture, Food and Forestry (D.M. 07/06/2012); SFP (Slow Food Presidia) are listed at [http://www.slowfoodfoundation.org](http://www.slowfoodfoundation.org)
Lactic acid bacteria fermentation induced a large modification of the profiles of immunoreactive protein of all the legume flours. A band of ca. 30 kDa, which was absent in the corresponding D, was found for LN, LU, LS, LP, and LA sourdoughs. The intensity of the 17 kDa band of fermented LV increased compared to D. The same signal became evident in all bean sourdoughs, especially for FL. Two protein bands were detected at ca. 30 kDa in FV, FSa, FBw and FBb. The same was found for CS and PS. Moreover, FV, FSa, FBw and FBb showed reactive bands containing the molecules with molecular mass lower than 30 kDa. In particular, the MTT assay was performed after treatment of Caco-2 cells with 0.1, 1, and 10 mg/ml of proteins for 24, 48, or 72 h.

Overall, all the WSE from D and S allowed a significant (P < 0.05) decrease of the cell proliferation compared to control (Fig. 2). Compared to the results obtained after 24 h-treatment, the inhibitory effect increased after prolonging the incubation to 48 h, and in many cases, to 72 h. Exceptions were the treatments with 10 mg/ml of proteins, which did not cause a further significant decrease of proliferation. The cytotoxic effect of WSE from S was in all the cases higher than the corresponding D. Besides the effect of the treatment duration, cytotoxicity increased proportionally to the concentration of proteins tested.

After 24 h of treatment (Fig. 2a), a weak anti-proliferative effect was found for WSE from D and S at 0.1 mg/ml of proteins. The strongest effect corresponded to LN: 72.25 ± 2.12 and 69.54 ± 1.88 % of vitality for D and S, respectively. When WSE from S were used at concentration of 1 mg/ml, the vitality of Caco-2 cells decreased by
Table 3 Microbiological analyses of the Italian legume flours

| Legume | Total mesophilic aerobic bacteria | Lactic acid bacteria | Yeasts | Molds | Total enterobacteria |
|--------|----------------------------------|---------------------|--------|-------|---------------------|
| FL     | 3.52 ± 0.12<sup>b</sup>         | 1.93 ± 0.12<sup>b</sup> | 2.77 ± 0.20<sup>c</sup> | 3.51 ± 0.13<sup>a</sup> | 0.82 ± 0.11<sup>b</sup> |
| FCo    | 3.60 ± 0.20<sup>b</sup>         | 2.49 ± 0.22<sup>a</sup> | 1.84 ± 0.20<sup>d</sup> | nf               | 0.53 ± 0.09<sup>f</sup> |
| FCu    | 3.23 ± 0.22<sup>b</sup>         | 2.54 ± 0.12<sup>c</sup> | 1.11 ± 0.05<sup>b</sup> | 3.33 ± 0.21<sup>a</sup> | 0.82 ± 0.11<sup>b</sup> |
| FS     | 4.21 ± 0.18<sup>a</sup>         | 2.23 ± 0.23<sup>a</sup> | 2.14 ± 0.21<sup>c</sup> | 3.31 ± 0.22<sup>c</sup> | 0.91 ± 0.07<sup>b</sup> |
| FV     | 2.92 ± 0.13<sup>c</sup>         | 1.14 ± 0.24<sup>c</sup> | nf    | 2.81 ± 0.15<sup>b</sup> | 0.51 ± 0.09<sup>f</sup> |
| Fsa    | 2.55 ± 0.25<sup>c</sup>         | 1.16 ± 0.11<sup>c</sup> | nf    | 2.47 ± 0.15<sup>c</sup> | 0.62 ± 0.13<sup>c</sup> |
| FBw    | 3.34 ± 0.19<sup>b</sup>         | 1.47 ± 0.15<sup>c</sup> | 1.30 ± 0.21<sup>c</sup> | 3.23 ± 0.16<sup>b</sup> | 0.73 ± 0.09<sup>f</sup> |
| Fbb    | 1.61 ± 0.32<sup>d</sup>         | 1.00 ± 0.16<sup>c</sup> | nf    | 2.45 ± 0.15<sup>c</sup> | 0.82 ± 0.09<sup>f</sup> |
| CM     | 2.84 ± 0.22<sup>c</sup>         | 1.31 ± 0.18<sup>c</sup> | 1.69 ± 0.23<sup>c</sup> | 2.47 ± 0.22<sup>c</sup> | 0.91 ± 0.14<sup>b</sup> |
| CV     | 2.47 ± 0.14<sup>c</sup>         | 1.47 ± 0.21<sup>c</sup> | 1.95 ± 0.09<sup>d</sup> | 2.30 ± 0.24<sup>c</sup> | 0.9 ± 0.09<sup>f</sup> |
| CS     | 2.25 ± 0.15<sup>c</sup>         | 1.69 ± 0.23<sup>c</sup> | nf    | 2.00 ± 0.21<sup>c</sup> | 0.8 ± 0.10<sup>d</sup> |
| CC     | 2.32 ± 0.23<sup>c</sup>         | 1.84 ± 0.22<sup>c</sup> | 4.23 ± 0.22<sup>a</sup> | nf               | 0.53 ± 0.05<sup>f</sup> |
| LN     | 2.69 ± 0.21<sup>c</sup>         | 1.77 ± 0.24<sup>c</sup> | 1.47 ± 0.13<sup>d</sup> | 2.04 ± 0.23<sup>c</sup> | 1.32 ± 0.10<sup>d</sup> |
| LU     | 2.36 ± 0.24<sup>c</sup>         | 1.84 ± 0.25<sup>c</sup> | 2.08 ± 0.15<sup>c</sup> | 2.30 ± 0.20<sup>c</sup> | 0.64 ± 0.11<sup>c</sup> |
| LS     | 2.95 ± 0.09<sup>c</sup>         | 1.47 ± 0.24<sup>c</sup> | 2.11 ± 0.15<sup>c</sup> | nf               | 0.75 ± 0.13<sup>b</sup> |
| LP     | 4.31 ± 0.11<sup>a</sup>         | 1.30 ± 0.25<sup>c</sup> | nf    | 3.23 ± 0.20<sup>d</sup> | 0.52 ± 0.22<sup>d</sup> |
| LA     | 3.69 ± 0.16<sup>b</sup>         | 1.30 ± 0.08<sup>c</sup> | 3.50 ± 0.15<sup>b</sup> | 2.11 ± 0.20<sup>c</sup> | 0.72 ± 0.14<sup>b</sup> |
| LV     | 2.61 ± 0.17<sup>c</sup>         | 1.00 ± 0.08<sup>c</sup> | 2.47 ± 0.12<sup>c</sup> | 2.23 ± 0.15<sup>c</sup> | 0.82 ± 0.12<sup>b</sup> |
| FS     | 2.47 ± 0.23<sup>c</sup>         | 1.30 ± 0.10<sup>c</sup> | 1.30 ± 0.21<sup>c</sup> | nf               | 0.92 ± 0.13<sup>c</sup> |

Total mesophilic aerobic bacteria were estimated on Plate Count Agar (PCA), lactic acid bacteria on agar MRS; yeasts and molds on Yeast extract Peptone Dextrose Agar (YPD-y and YPD-m, respectively); and total enterobacteria on Violet Red Bile Glucose Agar (VRBGA). Details are reported in “Methods” section. The data are the means of three independent experiments ± standard deviations (n = 3).

*** Values in the same column with different superscript letters differ significantly (P < 0.05).

ca. 10%. In particular, WSE from CC sourdough caused the highest decrease of vitality (58.47 ± 2.08%). At the highest concentration assayed, the most remarkable cytotoxicity was found for CC, LN, and PS sourdoughs (30.49 ± 1.13, 36.97 ± 1.01, and 35.13 ± 0.95%, respectively). The same trend was found after 48 h of treatment (Fig. 2b), although lower values of vitality were observed. The lowest proliferation was found for CC sourdough, when treatments were done with 1 mg/ml (39.59 ± 1.05%), and for CC, LN, and PS sourdoughs, when 10 mg/ml of proteins (30.67 ± 0.87, 29.32 ± 1.11, and 37.75 ± 1.23%, respectively) were used.

After 72 h of treatment (Fig. 2c), the vitality of Caco-2 cells was lower than that found after 24 or 48 h for treatments performed with 0.1 and 1 mg/ml of proteins, while the use of 10 mg/ml of proteins was less effective than 48 h-treatments, with the only exception of FL sourdough.

Transcriptional regulation of filaggrin (FLG) and involucrin (IVL)

Human colon adenocarcinoma cells (Caco-2) were treated with the partially purified and freeze-dried WSE from legume D and S aiming at investigating filaggrin (FLG) and involucrin (IVL) gene expressions through RT-PCR. Treatments were lasting 4, 8, and 24 h. After 4 h of exposure (Fig. 3A), a significant (P < 0.05%) over-expression of FLG was found only for fermented PS. All the other WSE, both from D and S, did not cause significant (P < 0.05) variations compared to LPS (positive control). After 8 h of exposure, a significant (P < 0.05) up-regulation of the FLG gene was also found for FL dough and FL, CV, and PS sourdoughs (Fig. 3B). The same trend was found for treatment lasting 24 h (Fig. 3C) with FL, CV, CC, and PS sourdoughs.

Regarding IVL gene, none of the extracts, except for PS sourdough, induced an expression after 4 h of treatment higher than that of LPS after 4 h of treatment (Fig. 4A). After 8 h of treatment, WSE extracted from S caused significant (P < 0.05) increases of the expression compared to the corresponding D, especially FL, CV, and PS (Fig. 4B). A longer treatment with the WSE (24 h) did not favour a further increase of the gene expression. Only PS sourdough caused the same expression of IVL gene under all the conditions assayed (Fig. 4C).

Identification of lunasin-like polypeptides

Based on the results from the MTT assay on Caco-2 cell, the immunoreactive protein bands of FL, CV, and PS sourdoughs were recovered from Tris-Tricine gels, and subjected to tryptic digestion, and HPLC coupled to nanoESI-MS/MS analysis.
Ten different proteins were identified through the Mascot research on the NCBI database (Table 5). It can be hypothesized that lunasin-like polypeptides were released from native legume proteins via proteolysis during sourdough fermentation. This should explain the lack of correspondence between the molecular masses of the proteins identified and the polypeptide bands revealed by western blot analyses. The immunodetected polypeptide bands from FL sourdough corresponded to the following matches: Subtilisin inhibitor 1 (P16064), Legumin A2 (P15838) and Phaseolin (P02853). The protein bands from PS sourdough were identified following matches: Subtilisin inhibitor 1 (P16064), Legumin A2 (P02858), legucoagglutinating phytohemagglutinin (P05087), phatogenesis related protein (CAAA56142) and seed linoleate 9S lipoxygenase-3 (P09918). The proteins from PS sourdough were identified as: Proviciolin (P02855), seed linoate 9S-lipoxygenase-2 (P14856), seed biotin-containing protein SBP65 (Q41060) and Albumin-1 C (P62928). All the database matches corresponded to proteins previously identified from legumes. For five proteins, the exact correspondence between the species analyzed and the database matches was not found, especially for CV. Probably, this was due to the limited number of legume protein sequences previously identified and included in the NCBI database.

The protein sequences were compared to soy lunasin sequence (deposited at the National Center for Biotechnology Information, NCBI database with the accession number AAP62458) using BLAST. Except for Proviciolin identified that was identified in PS sourdough, all the sequences might be aligned with different lunasin epitopes (Fig. 5). The longest alignments were found for Legumin A2, corresponding to fragments (f) 24–38 and 33–41 of the soy lunasin sequence, and Phaseolin (f13–28) from FL sourdough; pathogenesis related protein and seed-linoate 9S lipoxygenase-3 (f9–32 and f1–18, respectively) from PS. The alignments showed identities from 26 to 67 % (pathogenesis related protein and subtilisin inhibitor1/leucoagglutinating phytohemagglutinin, respectively) and positives from 35 to 93 % (Albumin-1C and Legumin A2, respectively).

#### Peptidase activities

The peptidase activities of the two lactic acid bacteria strains used as starters were assayed using relatively specific synthetic substrates. Significant ($P < 0.05$) differences were found between strains. Compared to that
Fig. 2 Effect of lunasin-like polypeptides on the Caco-2 cells proliferation. Box-plot showing aggregate data for human colon adenocarcinoma (Caco-2) cells proliferation after treatments of 24 (panel a), 48 (panel b), and 72 h (panel c) with water/salt soluble extracts obtained from control doughs (D), without bacterial inoculum, and sourdoughs (S), started with selected lactic acid bacteria, made with Italian legume flours. Protein concentration of 0.1, 1.0, and 10 mg/ml were assayed. Data were expressed as the mean percentage of viable cells compared to the control culture, grown in basal media without the addition of the WSE. The centre line of the box represents the median (open square), the top and bottom of the box represent the 75th and 25th percentile of the data, respectively. Outliers (open circle) and extremes (asterisk) are represented.

of *L. plantarum* C48 (5.87 ± 0.02 and 3.52 ± 0.02 U), PepN activity on Leu-p-NA and endopeptidase activity (PepO) of *L. brevis* AM7 were higher (6.83 ± 0.08 U and 4.31 ± 0.03). Also PepA activity was higher in *L. brevis* AM7 (14.02 ± 0.04 U). That of *L. plantarum* C48 resulted 35 % lower. This latter strain showed higher tripeptidase (PepT) activity (9.12 ± 0.02 vs. 5.61 ± 0.05 U). No significant differences (*P* > 0.05) were found between the two strains for PepT (7.20 ± 0.04 and 7.12 ± 0.05 U) and PepX (0.95 ± 0.03 and 1.01 ± 0.02 U) activities.

Discussion

Despite the beneficial effects on the human diet, the worldwide consumption of legumes is declining [15] and below the recommended dose [16]. One potential way to increase the consumption of legumes could be the rediscovery of traditional and local varieties, and, especially, the use of legumes into novel and healthy foods, also exploiting the potential of non-conventional processing [17]. The complementation between cereal and legume flours into new formulas may deserve an interest either to increase the levels of biogenic compounds or to fulfill nutritional deficiencies of cereal-based diets [18].

Traditional Italian legumes, all with product certifications and belonging to *Phaseolus vulgaris*, *Cicer arietinum*, *Lathyrus sativus*, *Lens culinaris* and *Pisum sativum* species, were used in this study. Seeds were milled, and flours were subjected to sourdough fermentation, using
the selected *Lactobacillus plantarum* C48 and *Lactoba-
cillus brevis* AM7.

Compared to cereals, all legume flours showed elevated amount of protein (mainly grass pea and several lentil flours), FAA (>2 g/kg), dietary fibers, and ash. In particular, grass pea varieties had the highest concentration of pro-
teins, while pea, chickpea, and grass pea flours contained the highest levels of dietary fibre. The microbiota of leg-
ume flours was poorly represented by lactic acid bacteria and yeasts. All the data agreed with previous findings [19].

Recently, the potential of sourdough fermentation was exploited to enhance the nutritional and functional fea-
tures of legume flours [19]. Apart from the legume species and variety, sourdough fermentation with selected starters is a suitable biotechnology option either to increase the nutritional and functional value or to decrease the levels of anti-nutritional factors [19]. Selected sourdough lactic acid bacteria were able to decrease the concentra-
tion of raffinose up to ca. 64 % and a similar trend was found for the concentration of condensed tannins [19].

The sourdough fermentation also increased the concen-
tration of GABA, and promoted antioxidant and phytase activities compared to the raw flours [19].

According to protocols used for cereal sourdough fer-
mentation [20], legume flours were started with *Lactoba-
cillus brevis* AM7 and *Lactobacillus plantarum* C48 [21].

As previously shown [19], the conditions of incubation (24 h at 30 °C) allowed the optimal growth and metabo-
lism of selected lactic acid bacteria [19, 22]. As expected, fermentation caused a marked decrease of the values of pH as well as an increase of TTA and of the concentra-
tion of peptides and FAA, especially when kidney bean flours were used.

Based on their proteolytic activity towards vegetable proteins [20], the use of sourdough lactic acid bacteria for synthesizing bioactive peptides deserves a marked interest [23]. Bioactive peptides derived from food pro-
teins may possess physiological properties beyond the role in nutrition. These properties are influenced by the protein source, enzyme and processing conditions used [24]. Most of the research related to bioactive peptides and cancer was focused on lunasin [25] and, recently, on pulse hydrolysates [2].

To the best of our knowledge, no literature data dealt with the presence of lunasin or lunasin-like polypeptides from native protein sequences of Italian pulses, and with the effect of sourdough fermentation on bioactivity and bioavailability.

Proteinase activity and a large portfolio of peptidases are the pre-requisites to liberate bioactive peptides from native oligopeptides [7, 26]. Although with some differ-
ces, the two starters used showed different peptidase activities.

Western blot analyses showed that the sequence of lunasin sequence was absent in all the legume flours, and, therefore, in the corresponding sourdoughs. Never-
evertheless, immunoreactive polypeptides with molecular masses lower than 30 kDa were detectable in all the samples. In some cases, immunoreactive polypeptides appeared as multiple bands in western blot, probably due to the presence of multiple fragments differing for few aminoacid residues [12]. Regarding unfermented doughs, proteins with high intensity signals were found for all the bean, followed by lentil varieties. After lactic acid bacteria fermentation, the number and the intensity of the proteins reacting with the anti-lunasin antibody increased for all the legume flours. This was probably due to an acid activation of endogenous proteinases, respon-
sible for primary proteolysis, and peptidase activities by lactic acid bacteria, which completed the hydrolysis (sec-
ondary proteolysis). All the reactive protein bands had molecular masses higher than that of lunasin. Although some differences were found among varieties, one repre-
sentative was chosen for each legume species to assay the effect on Caco-2 cells proliferation. Caco-2 cells derived from a colonic tumor and have a cancerous phenotype, and can be cultivated to become confluent. In this case, they differentiate into enterocyte-like cells [27].
A

IVL mRNA (relative quantity chart)

LPS  FL  CV  CC  LN  PS

B

IVL mRNA (relative quantity chart)

LPS  FL  CV  CC  LN  PS

C

IVL mRNA (relative quantity chart)

LPS  FL  CV  CC  LN  PS

Legend:

D
S
| WSE | Protein | Sequence<sup>a</sup> | NCBI accession number | Theoretical mass (kDa) | Sequence coverage | Mascot score |
|-----|---------|----------------------|-----------------------|-----------------------|------------------|--------------|
| FL-S | Subtilisin inhibitor 1 (Vigna angularis) | QEQGTNPSEQMVLPLRNYKQALETNTPTKTSWPLGVTVAEQAETKIQEEMVDAIQVSPHSDFVTADYNPKRVRVLSVDESNKVTTPSGLYNAPQIEFOQQNGYFGMVPGPCTFEPEEQSEDQECGRRYRDRIRQKVNREGRIDAVPTGMFWMNQDPTVPAVSLIDRIRSNLOLDMPHRFYLANGNHQEFELRYOHQOQQGKOEOEENGNNIFSFFKRDPLLEDAFVVNRHRIVLRQLGRNDEEDEEGAKLKVXLGSISPEKXGARHGRSROEEDDEDEEDEPQROPIVGRSGRQSEEDEEERQPOTRRRGGSSIDKERRGSGQKSRQGQDNQLEETVCTAIXLRLNGPSSPDYNEPAEGRKTVTSIADNLRLWRLKSLASEKIHGUNKAMFVPHYNLNSIYALKGRARLOVVNCOQNTVDGELEAGALTVPONYAYAAKSLSRFYSYAFKNTDRAGIALGTSSVINLPLDIVAATFDNQLQNEARQLKSNNPFFLVPARQSENARAL | P16064 | 10.38 | 82 | 114 |
| Legumin A2 (Pisum sativum) | MATXKLLASLSCFLLLGLGCFAFLRQPEQOEQNECQLERLNALPEPDNRIRSEGGLUTWNPNNQQFLRCAGIALSATLQLHLNALLPRPYYSNAPQIEFOQQNGYFGMVPGPCTFEPEEQSEDQECGRRYRDRIRQKVNREGRIDAVPTGMFWMNQDPTVPAVSLIDRIRSNLOLDMPHRFYLANGNHQEFELRYOHQOQQGKOEOEENGNNIFSFFKRDPLLEDAFVVNRHRIVLRQLGRNDEEDEEGAKLKVXLGSISPEKXGARHGRSROEEDDEDEEDEPQROPIVGRSGRQSEEDEEERQPOTRRRGGSSIDKERRGSGQKSRQGQDNQLEETVCTAIXLRLNGPSSPDYNEPAEGRKTVTSIADNLRLWRLKSLASEKIHGUNKAMFVPHYNLNSIYALKGRARLOVVNCOQNTVDGELEAGALTVPONYAYAAKSLSRFYSYAFKNTDRAGIALGTSSVINLPLDIVAATFDNQLQNEARQLKSNNPFFLVPARQSENARAL | P15838 | 59.63 | 62 | 410 |
| Phaseolin (Phaseolus vulgaris) | MRRARVPLLGLFLALISASAFSATSLLGEQEQDONQNFYFNSDNSWNTLKNQYGHPRQLQFQDQSKRLQNLLEDYRLPEFKPETELLPOQADAEILVQRSGAILVWKPDPDREYFLTSDNPSHDQKIPAGIFLYLNPDPKDELIRLLAMPSNQPIHEFFLSSTEAQOSQYLGESFSKHEILSFNSKFEINRLVEEEQEGQWGNIDSEOKLELHAKSSRSLSKQODINTNGEFQNTERTDNSLWLISSIEEMGALFYPHYSSAMVILVNEGEAMVLEVPKGMNKTLEYSEYSAELKDDIVJDPAPYVAIAKATSWNKTFTGGINANNNIRNLAGKTDNMSJICRALKDOKDVLGLTSGGDELETEVMKINKOSQSYFDVAAAHHQQEFOQKGRKGFVYY | P02853 | 47.540 | 75 | 515 |
| CV-S | Leucoagglutinating phytohemagglutinin (Phaseolus vulgaris) | MASSKFFTVLFVTLTHANNSENDIFNFQRFNTERNLQRDASVSSGGQLRLNNGPVGSLGRANSAPIQWDNTTGTAVASFSTFTFNQVNPINAGPDQLFAALVPVSQOKDGFLGDGSNSNHFTVAVEFDSTYLNKILWDPETERHDSNPRSTTRFDWPVNGNEAEVLTIVDSTNLELUASLVPVOSKFTSDVNDVLSLPEWSVGPSSATGINKGVTENVLSWFFSALKSLDGTTSEGPNLNLVNLNKL | P05087 | 29.54 | 61 | 506 |
| Pathogenesis related protein (Cicer arietinum) | MQVTFTEQESTTVPPAKLYKAMIKDAVIIKPKDAKJTKUVTPVECGNGGPTIKLKFVEQGQTLYVLHKEIADEALNGYNYSVCGAGLSETVERYHEAKLCEQPGNGSISOKVSVKYQKTGDAPKPNEKEVQEGKAKGDLIFKAIEGYVLANPYN | CAA56142 | 16.93 | 83 | 408 |
### Table 5 continued

| WSE       | Protein                        | Sequence<sup>a</sup>                                                                 | NCBI accession number | Theoretical mass (kDa) | Sequence coverage | Mascot score |
|-----------|--------------------------------|--------------------------------------------------------------------------------------|-----------------------|-----------------------|-------------------|-------------|
|           | Seed linoleate 9S-lipoxygenase-3  | MFSVGTVGILNNGRHRKGTVILMRKNNVLDINSITTVGVVIGQFIDLGSTYDNLFAFLGQVSLQQUESATKPDATKGGKIKETFLGLQISSLPTLGAQOAFKIHFEWDDDMGPGAFYKKNMOTEFDFLSILTLIDIDPNHGSIYVLCNSWYNAKHIDRFFANQTYLPSETPAPILVHYREEELNLGRGDGTGKEXERIYDYVYNLDGNPDSGENHARPVLQGGEPTYTPPRGRTGKPRTPKDPNSERS5DYVLPRLDEAFGHKSSDFFTYGLKAVSQNVVPAPESYFDFDLNTPFTEDSVDFDLVGEYGGKPTNILSQSPLPYVLBERTDQENGLEYPPKXVQ5/SRGSWMTDEFAREM | P09918 | 97.57 | 43 | 207 |
| PS-S      | Provicilin (Pisum sativum)      | DNAEIKLLEHEKETHHRRLGRDKQRQSQEKMVWYSSKQIEEELKNAKSS5KSSVSESEFPLKSSDDPPLYSQYKKEEITPKKNPQLQDLDFNPYVEKEGSLSWLPHYNSRMAMVTNEGKGD | P02855 | 31.52 | 53 | 589 |
**Table 5 continued**

| WSE Protein Sequence | NCBI accession number | Theoretical mass (kDa) | Sequence coverage | Mascot score |
|----------------------|-----------------------|------------------------|------------------|--------------|
| Seed linoleate 9S-lipoxygenase-2 (Pisum sativum) MFPNVTGLNKGHKRIQTVGVLMLRNAFLDNTIVSKGGGVHKVSDGI NIKGTLDGLTAFLGRSVESQCVASATKSDANHGKKGKVGDVFLEGV LSLFPLTLGASEAFNHFDWHDHMGP121GAFYKMNQMEVEF FLKSLTLEDVPNHTFIRCVSNWVNSKLYKSPRIFFANKLYLPSET SPSLVKRYEELQLOLDGDTGERKLRHLYDYDVNDLNPWDE GEHLRIPLGGSHFHPYPRGRTGRTFRKDFSNKPATTETVPPRDEN FGHKLSDFLAYKSVSQCVPAPSFALDNFTPNEFDSQDVVRNF LFEGKIKILDLVATLPSLPAKEIFRTDQEGQUTFKPHHRVYVSANW MTDDEERELMLAGVYNPCMIRLQFEFPPKSNLPEVDGHTKSVSIVD VLNLDGCCIDEALASGRFLDVLGHTIPFLRRINETSAYKATRITLFLK ENGTLKIPAEISLPHPDGRKVJFSGVSKLPADEGVESTWILLAAYV WNDSCYHQOMSHWLTHAVIEPVIAHNLQSVVHPINKLAPAHY RDTMNNINALARDULANLQERFSFLPSKYYAEMSSAYKYVWF DTQALPNLIDKRNAYWDESSEYGLBKLEEPYVADQLGELWTAIKT WQDYVSLSYATNDDIKNSLQOMWKEVKGQLDKPWWQP KLOTDELEVCCTIIWTASLAHAVNFQGYPYGLILNRPLTSRRLL PEIGFAEYDEMIVSKQKAYLRTTPFQTLDDLSSRLASHDARGYL QORENPMYTSKALQAFQKFNKLAEEIKLNNNDPSLHYHRVG PVQNPYTLHPSSKQGLTFRPGPNSISI |
| Seed biotin-containing protein SBP65 (Pisum sativum) MASEQSLRRENTERTKQNAEVDVPVF.TestCaseLGPQNFQPLQEAOVLIRGKAPLANSYSEYLDRARVPLNANP EHRVREKEDFQGVRMDQMGKQMESKGGNKLAEEDRLTDTSRM VGTGPHKEASGKGQWEERERAREAMEEEEKLTMEEISKYRN QAQEQSALEALSEEAQERERKOSNATIgLNPQNTAQKMQGSHAEQAKQ DATFNEKQOGYEMGTENVNPATSEQARATNQRTLQDGQYET ARTDTVSNARTAECYKAEAVQAKDVTLEGTKTAAEKCAAC AAEBACKVAVDILKELKATAVGWTATHYQLTQDLQDTRAANNAE VGAVYVAKAESLAKSVETKGLASAGETKAFYRTKEESESWE YEAKRASQLOEEGEEILPSGKKILPSERTQAQTNKREQKVGQG KSGLDGAVETYSDQGSMKPDADKNTKVEHGGITITPFXPQGDAQ GYLDIAETIAAEHYYYTVVVEDDEEVEKSMQNKGDHSLDRAKLY EGYRAPKNIVS |
| Albumin-1 C (Pisum sativum) MASWKLSLFLAFATMMLTKNVAGCNGCVSFPEDEPPCGSPLCRCI PAGLIVGNRNPYGFVRNLTNDEFPHRNCLSEDADCRKKSXSGTFCHGYP NPDEFYCGFCASKSAEADFVSKTPKDLLKSVSTA |

Identification was carried out by nano-LC-ESI–MS–MS. Polypeptides were purified from water/salt-soluble extracts obtained from sourdoughs (S) made with Fagiolo di Lamon (FL), Cece dell’Alta Valle del Misa (CV), and Pisello riccio di Sannicola (PS).

* Single-letter amino acid code is used
Polypeptides able to regulate cell proliferation and survival were already liberated by enzyme hydrolysis of plant and animal proteins [28]. Such peptides, inhibiting cell growth or promoting apoptosis, could have protective effects on tumor growth at the digestive tract level [29, 30]. In particular, it was found that the antitumoral mechanism of soybean lunasin is related to the capacity of the peptide to bind the deacetylated histones, causing the acetylation inhibition [31, 32]. Girón-Calle et al. [28] assessed the cancer cell proliferation after treatment with chickpea hydrolysates made with pepsin/pancreatin. These hydrolysates inhibited the proliferation of human epithelial colorectal adenocarcinoma cells (Caco-2) and monocytics leukemia cells (THP-1) up to 48 and 78 %, respectively. Hydrolysates from the common bean (P. vulgaris) varieties Negro 8025 and Pinto Durango inhibited the inflammation in lipopolysaccharide-induced macrophages through suppression of NF-κB pathways [2]. After hydrolysis with Alcalase, proteins extracted from both the above varieties inhibited various markers of inflammation (cyclooxygenase-2 expression, prostaglandin E2 production, inducible nitric oxide synthase expression, and nitric oxide production) [2]. Inflammation and cancer are linked, chronic inflammation predisposes individuals to various types of cancer and inflammatory mediators and cells are involved in the migration, invasion, and metastasis of malignant cells [33]. The suppression of pro-inflammatory pathways may provide opportunities for both prevention and treatment of cancer [34].

In this study, extracts from legume doughs regulated the proliferation of Caco-2 cells, including those obtained from control doughs, in which very weak signals of immunoreactive bands were found with the western blot analysis. It can be hypothesized that the effect could be related to the contribution of some legume proteins or peptides able to act as anticancer compounds in their native form, causing cytotoxicity and apoptosis of tumoral cells [2, 5]. Nevertheless, a strong inhibition of the Caco-2 cell proliferation was found only after lactic acid fermentation. A marked inhibition was found when cells were treated for 24 h with the fermented legume extracts at concentrations from 1 to 10 mg/ml. Sourdoughs from FL, CV, and PS flours were the most active, showing a decrease of Caco-2 cells vitality up to 70 %. A correlation between the anti-proliferative effect of protein hydrolysates towards Caco-2 cells and the in vivo growth inhibition of tumors in the digestive tract was demonstrated [29, 30].

The nanoESI-MS/MS spectrometer analysis of the immunoreactive protein bands from water/salt-soluble extracts of FL, CV and PS sourdoughs allowed the identification of ten legume proteins. According to the molecular masses of the identified proteins, it can be hypothesized that immunoreactive polypeptides are encrypted into the native sequences and released as fragments during lactic acid bacteria fermentation. Nine of them showed similarities to soy lunasin sequence, probably related to their recognition by the anti-lunasin antibody.

Using Caco-2 cells, the capacity to induce the expression of human FLG and IVL genes by sourdough fermented legumes was investigated. FLG and IVL are important proteins for the formation of the epidermal skin barrier [35, 36]. FLG aggregates keratin filaments and provides a cytoskeleton for the cornified envelope [35]. The expression of FLG was markedly induced by PS sourdough, and under some of the assayed conditions (e.g., 8 h of treatment), also by CV and FL sourdoughs. Compared to the unfermented dough, the level of IVL gene expression markedly increased when Caco-2 cells were subjected to 24-h treatment with FL, CV, and PS sourdoughs. In particular, the sourdough made with PS was effective under all the conditions likely it was observed for FLG gene expression. IVL serves as a substrate for the covalent attachment of ceramides to the cornified envelope [35]. An improvement of the IVL activity may result in an increase of the ceramide binding activity, which, in turn, positively affects the barrier function [37, 38].
Conclusions
Nowadays, an unexpected and considerable number of small proteins and peptides are available as drugs [39]. They show high bioactivity, target specificity and wide spectrum of therapeutic actions, with low levels of toxicity, structural diversity, and absence or low levels of accumulation in body tissues. However, the manufacturing protocols (e.g. chemical synthesis and recombinant transgenic approach) are very expensive, representing a hindrance for the use as therapeutic peptides [40]. The alternative option is the exploitation of the potential of bioactive peptides derived from food proteins. Legume flours were already proposed as adjuvant ingredients into a range of baked products and snacks for increasing the nutritional value [41–44], as well as sourdough fermentation was shown to have a functional potential [19, 45, 46]. Despite the need to have an in vivo confirmation, the presence of lunasin-like polypeptides allows to hypothesize the use of sourdough fermented legumes in pharmaceuticals preparations (e.g., capsules and powders), protein hydrolysates or as purified peptide mixtures to be incorporated as health-enhancing ingredients in novel functional foods (e.g., leavened baked goods).

Methods
Legumes
Nineteen Italian legume varieties, belonging to the species Phaseolus vulgaris (Fagiolo di Lamon, Fagiolo di Controne, Fagiolo di Cuneo, Fagiolo Stregoni, Fagiolo Vellutina, Fagiolo di Saluggia, Fagiolo Badda di Polizzi—white, and Fagiolo Badda di Polizzi—black), Cicer arietinum (Cece di Merella and Cece dell’Alta Valle del Misa), Lathyrus sativus (Cicerchia di Serra de Conti and Cicerchia di Campodimele), Lens culinaris (Lentichcia di Castelluccio di Norcia, Lentichcia di Ustica, Lentichcia di Santo Stefano di Sessanio, Lentichcia rossa di Pantelleria, Lentichcia di Altamura and Lentichcia di Villalba), and Pisum sativum (Pisello riccio di Sanni cola), were used in this study. All legumes were chosen among the Italian pulses that have specific product certification (Table 1). Flours were obtained from whole legume seeds through the laboratory mill Ika-Werke M20 (GMBH, and Co. KG, Staufen, Germany). Protein (total nitrogen ×5.7), ash and moisture contents were determined according to AACC approved methods 46-11A, 08-01 and 44-15A, respectively [47]. Lipids were determined by Soxhlet method. Total carbohydrates were calculated as the difference, using the following formula: [100 – (proteins + lipids + ash + moisture)]. Proteins, lipids, carbohydrates and ash were expressed as % of dry matter (d.m.). The determination of total dietary fiber was carried out by the enzymatic–gravimetric procedure approved by the Association of Official Analytical Chemists [48], as described by Lee et al. [49].

Microbiological analyses of the flours
Ten grams of flour were homogenized with 90 ml of sterile peptone water [1 % (wt/vol) of peptone and 0.9 % (wt/vol) of NaCl] solution. Lactic acid bacteria were enumerated using MRS (Oxoid, Basingstoke, Hampshire, UK) agar medium, supplemented with cycloheximide (0.1 g/l). Plates were incubated, under anaerobiosis (AnaeroGen and AnaeroJar, Oxoid), at 30 °C for 48 h. Cell density of yeasts and molds were estimated on Yeast Extract Peptone Dextrose Agar (YPD) (Oxoid) medium (pour and spread plate enumeration, respectively), supplemented with chloramphenicol (0.15 g/l), at 30 °C for 72 h. The attribution (yeasts/molds) was confirmed by microscope observation. Total mesophilic aerobic bacteria were determined on Plate Count Agar (PCA, Oxoid) at 30 °C for 48 h. and total enterobacteria were determined on Violet Red Bile Glucose Agar (VRBGA, Oxoid) at 37 °C for 24 h.

Fermentation
Sourdough lactic acid bacteria, belonging to the Culture Collection of the Department of Soil, Plant, and Food Sciences (University of Bari, IT) and previously selected for some biotechnological features, were used. Lactobacillus plantarum C48 showed the capacity of synthesizing relevant amount of γ-aminobutyric acid (GABA) [21], and Lactobacillus brevis AM7 was characterized by high proteolytic activities towards cereal and legume flours [12]. Starters were cultivated separately on MRS broth at 30 °C for 24 h. Cells were harvested by centrifugation (10,000 × g, 10 min, 4 °C) until the late exponential phase of growth was reached (ca. 10 h), washed twice in 50 mM sterile potassium phosphate buffer (pH 7.0) and re-suspended in tap water at the cell density of ca. 8.0 log cfu/ml. Each legume flour was mixed with tap water, containing the bacterial suspension (initial cell density of 7.0 log cfu/g of sourdough for each strain). Doughs, having dough yield (DY, dough weight ×100/flour weight) of 160 (corresponding to 62.5 and 37.5 % of flour and water, respectively), were mixed at 60 °C for 5 min with a IM 58 high-speed mixer (Mecnosud, Flumeri, Italy) and incubated at 30 °C for 24 h. The most common DY value for cereal-based sourdoughs (160) was applied [20, 22, 46, 50]. After fermentation, legume sourdoughs (S) were compared to control doughs (D), without bacterial inoculum, prepared as described above (DY 160) and incubated under the same conditions. S and D were stored at −20 °C before the chemical analyses, while microbiological analysis was carried out before freezing. All the
doughs were obtained in triplicate and each of them was analyzed twice.

**Characterization of fermented flours**

The values of pH were determined on-line by a pHmeter (Model 507, Crison, Milan, Italy) with a food penetration probe. Total titratable acidity (TTA) was determined after homogenization of 10 g of dough with 90 ml of distilled water, and expressed as the amount (ml) of 0.1 M NaOH needed to reach the value of pH of 8.3.

The water/salt-soluble extract (WSE) of D and S was prepared according to Weiss et al. [51], and used to analyze free amino acids (FAA) by a Biochrom 30 series Amino Acid Analyzer (Biochrom Ltd., Cambridge Science Park, UK) with a Na-cation-exchange column (20 by 0.46 cm internal diameter), as described by Rizzello et al. [50].

**Peptidase activities of the starters**

General aminopeptidase type N (EC 3.4.11.11; PepN), specific aminopeptidase type A (EC 3.4.11.7; PepA) and endopeptidase (EC 3.4.23; PepO) activities were determined as described by Gobbetti et al. [13], using, respectively, Leu-p-nitroanilides (p-NA), Glu-p-NA and NCBZ-Gly-Gly-Leu-p-NA (Sigma Aldrich Co.) as relatively specific substrates. The assay mixture contained 900 μl of 2.0 mM substrate in 0.05 M potassium phosphate buffer, pH 7.0, and 100 μl of cell suspension (9 log cfu/ml). The mixture was incubated at 30 °C for 1 h and the absorbance was measured at 410 nm. The data obtained were compared to standard curves set up by using p-nitroanilide [52]. One unit (U) of activity was defined as the amount of enzyme required to liberate 1 μmol/min of p-nitroanilide under the assay conditions. Tripeptidase (EC3.4.11.4; PepT), and X-prolyl dipeptidyl aminopeptidase (EC 3.4.14.5; PepX) activities were determined using Leu-Leu-Leu and Gly-Pro-Ala substrates (Sigma Aldrich Co.), respectively. Activities on tripeptides were determined by the Cd-ninhydrin method [52], [53]. The assay conditions were the same as those described for p-nitroanilide substrates. One unit (U) of activity was defined as the amount of enzyme required to liberate 1 μmol amino acid released per min under the assay conditions. The data obtained were compared to the standard curve set up by using leucine [52].

**Western blot analysis**

Protein concentration of samples was determined by the bicinchoninic acid method (Thermo Scientific, Rockford, IL, USA), using bovine serum albumin (BSA) as standard protein. SDS-PAGE was performed with samples and synthetic lunasin diluted in tricine sample buffer (Bio-Rad, Richmond, CA, USA), containing 2 % (v/v) β-mercaptoethanol, and heated at 100 °C for 5 min. Equal amounts of proteins (20–40 μg) were analyzed on Precast Criterion 16.5 % Tris-Tricine gels (Bio-Rad), and electrophoretic separations were carried out at 100 V, using Tris-Tricine-SDS as running buffer in the Criterion cell (Bio-Rad). For the attribution of the molecular masses, a Precision Plus Protein Standards mix (BioRad) was used. After SDS-PAGE separation, gels were soaked in transfer buffer (48 mM Tris, 39 mM glycine, 20 % methanol, pH 9.2) for 30 min. Proteins were electroblotted into nitrocellulose membranes by semidy transfer in a Trans-Blot SD (Bio-Rad) for 30 min at 18 V. Then, the membranes were blocked for 3 h in Tris buffered saline with 0.05 % (v/v) Tween 20 (TBST), containing 1 % (w/v) bovine serum albumin, (TBST-1 % BSA). Afterwards, the membrane was washed three times with TBST, and incubated overnight at 4 °C with lunasin polyclonal primary antibody (diluted 1:2000 in TBST-0.1 % BSA). After washing with TBST, the membrane was incubated overnight at 4 °C with horseradish peroxidase-conjugated mouse anti-rabbit secondary antibody (Santa Cruz Biotechnology, Dallas, TX, USA; 1:3000 in TBST-0.1 % BSA). Finally, the membranes were washed six times with TBST, and visualized by chemioluminescence using the detection agent Amersham TM ECL Prime (GE Healthcare, Chalfont St Giles, UK), according to the manufacturer’s recommendations. Image acquisition (exposure time 1–4 min) was performed using the VersaDoc Imaging System (Bio-Rad).

**MTT assay in Caco-2 cells**

In order to assess the effect of legume D and S on cell proliferation, the viability of colon adenocarcinoma Caco-2 cells was measured using the 3-(4,5-dimethyl-2-y)-2,5-diphenyltetrazolium bromide (MTT) method [54].

Colon adenocarcinoma Caco-2 cells (ICLC HTL97023), provided by the National Institute for Cancer Research of Genoa (Italy), were cultured in RPMI medium, supplemented with 10 % Fetal Bovine Serum (FBS), 1 % 2 mM L-glutamine, 1 % penicillin (10,000 U/ml)/streptomycin (10,000 μg/ml) mixture, 0.1 % gentamicin and 0.1 % β-mercaptoethanol, and maintained in 25 cm² culture flasks at 37 °C, 5 % CO₂. Every 2 days, confluent cultures were washed with PBS 1× (without Ca²⁺ and Mg²⁺), split 1:3–1:6 using Trypsin/EDTA, and seeded at 2–5 × 10⁴ cell/cm², 37 °C, and 5 % CO₂. For the assay, Caco-2 cells were harvested at 80 % confluence with trypsin/EDTA, seeded at the density of 5 × 10⁴ cells/ml into 96-well plates, and then incubated at 37 °C, 5 % CO₂ for 24 h. Before analysis, WSE obtained from legume D and S were partially purified by ultra-filtration, using Vivaspin (GE Healthcare) centrifugal filter units (cut-off 30 kDa), following the manufacturer’s instructions.
Cells were exposed to partially purified and freeze-dried WSE at the following concentrations: 0.1, 1.0, and 10.0 mg/ml of proteins. Each WSE was tested in duplicate. A control in basal medium, without addition of WSE, was used. Incubation was carried out for 24, 48 and 72 h. After incubation, the medium was removed and replaced by 100 µl/well of dime-mixture, 0.1 % gentamicin and 0.1 % β-mercaptoethanol and maintained in RPMI, containing 10 % FBS, 1 % 2 mM l-glutamine, 1 % penicillin (10,000 U/ml)/streptomycin (10,000 µg/ml) mixture, 0.1 % gentamicin and 0.1 % β-mercaptoethanol. Then, the medium was removed and 100 µl/well of dimethyl sulphoxide (DMSO) were added to dissolve the purple formazan product. Plates were shacked for 15 min at room temperature and the absorbance was read at a wavelength of 570 nm, with a Biotek microplate reader (BioTek Instruments Inc., Bad Friedrichshall, Germany), and elaborated with the ELX808 software (BioTek Instruments Inc., Bad Friedrichshall, Germany). Data were expressed as the mean percentage of viable cells compared to control culture, grown in basal media without addition of WSE.

Identification of the lunasin-like polypeptides
After Western blot analysis, the immunoreactive bands, separated by Tris-Tricine SDS PAGE, were cut and stored in 20 % ethanol before identification. Protein identification was performed by Proteome Factory (Proteome Factory AG, Berlin, Germany). Protein bands were digested by trypsin (Promega, Mannheim, Germany) and analyzed by Nano-Liquid Chromatography-Electrospray Ionisation-Mass Spectrometry (nanoLC-ESI–MS/MS). The LC–MS system consisted of an Agilent 1100 nanoHPLC system (Agilent, Waldbronn, Germany), using a linear gradient from 2.5 % acetonitrile/0.5 % formic acid, then separated on a Zorbax 300SB-C18, 75 μm × 150 mm column (Agilent), using a linear gradient from 10 to 32 % B (solvent A: 5 % acetonitrile in water, solvent B: acetonitrile, both with 0.1 % formic acid). Ions of interest were data-dependently subjected to MS/MS, according to the expected charge state distribution of peptide ions. Proteins were identified by database search against the plant sequences of the National Center for Biotechnology Information, (NCBI, Bethesda, USA) protein database, using MS/MS ion search of the Mascot search engine (Matrix Science, London, UK). Only peptide matches with a score of 20 or above were accepted. The sequences of the protein identified were aligned using the BLAST on-line tools (http://blast.ncbi.nlm.nih.gov).

Transcriptional regulation of flaggrin (FLG) and involucrin (IVL) genes
Colon adenocarcinoma Caco-2 cells (ICLC HTL97023) were cultured in RPMI medium supplemented with 10 % FBS, 1 % 2 mM l-glutamine, 1 % penicillin (10,000 U/ml)/streptomycin (10,000 µg/ml) mixture, 0.1 % gentamicin and 0.1 % β-mercaptoethanol and maintained in 25 cm² culture flasks at 37 °C, 5 % CO₂. Caco-2 cells were incubated in 25-cm² culture flasks at 37 °C, under 5 % CO₂ atmosphere [55].

Subconfluent monolayers of Caco-2 cells were subjected to treatment with basal medium, containing 2.5 % FBS. The freeze-dried WSE, partially purified by ultra-filtration using Vivaspin (GE Healthcare) centrifugal filter units (cut-off 30 kDa), were dissolved in RPMI medium, at the concentration of 10 mg/ml and added to the culture media at the final concentration of 1 mg/ml. The control was the basal medium, containing 2.5 % FBS. Plates were incubated at 37 °C for 24 h, under 5 % CO₂. Samples were taken after 4, 8, and 24 h of treatment. Each experiment was carried out at least twice in triplicate. For quantitative real-time PCR (RT-PCR), total RNA was extracted from Caco-2 cells using the RiboSpin Minikit-GeneAll kit. cDNA was synthesized from 2 μg RNA template in a 20-μl reaction volume, using the Prime Script RT reagent kit (perfect Real time) (Takara). Total RNA solution (10 μl) was added to the Master Mix and subjected to reverse transcription in a thermal cycler (Stratagene Mx3000P Real-time PCR System, Agilent Technologies Italia Spa, Milan, Italy). The conditions were 37 °C for 15 min, 85 °C for 5 s, holding the samples at 25 °C.

The cDNA was amplified and detected using the same instrument and the TaqMan assay (Applied Biosystems). The following Taqman gene expression assays were used: FLG Hs00863478_g1 (FLG), IVL Hs00846307_s1 (IVL), and GAPDH Hs99999905_m1 (human glyceraldehyde-3-phosphate dehydrogenase, GAPDH). Human GAPDH was used as the housekeeping gene. PCR amplifications were carried out using 40 ng cDNA in a total volume of 20 μl. The reaction mixture contained 10 μL of Premix Ex Taq, 0.4 μl of RoxTM reference dye II, 1 μL of 20X TaqMan Gene Expression assay and 4 μl of cDNA.

PCR conditions were 95 °C for 30 s (for AmpliTaq activation), followed by 40 amplification cycles (95 °C for 5 s, 60 °C for 20 s). Analyses were carried out in triplicate. Based on preliminary results, the expression of FLG and IVL genes was also assayed by treating Caco-2 cells with lipopolysaccharide (LPS; Sigma Aldrich Co.) at 10 μg/ml in basal medium, containing 2.5 % FBS. Analyses were
carried out in triplicate. The average value of target gene was normalized using GAPDH gene and the values were elaborated automatically by the MXPro v.4.01 Stratagene software [56].

Statistical analysis

Data were subjected to one-way ANOVA; pair-comparison of treatment means was achieved by Tukey’s procedure at $P < 0.05$, using the statistical software, Statistica 7.0 for Windows. Student’s $t$ test was used for MTT assay (GraphPAD 6.0 for Windows).

Abbreviations

BSA: bovine serum albumin; D: dough; DY: dough yield; FBS: fetal bovine serum; FLG: filaggrin; GABA: y-aminobutyric acid; iLvl: involucrin; MT: 3-(4,5-dimethyl-2-yl)-2,5-diphenyltetrazolium bromide; S: sourdough; TBS: tris buffered saline; TTA: total titratable acidity; WSE: water/salt-soluble extract.

Authors’ contributions

CGR was responsible for the experimental design of the work, the identification of polypeptides, and the elaboration of all the data; BHL and SFT performed immunological analyses and purification of polypeptides; JAC carried out fermentations and microbiological analyses; DP and BN performed all the analyses on Caco-2 cells; RC performed the biochemical characterization of samples; MG was the supervisor and the coordinator of the research unit. All authors read and approved the final manuscript.

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Competing interests

The authors declare that they have no competing interests.

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