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***Proteomic analysis of the metabolic fractions
in modern and old wheat genotypes: a
qualitative and quantitative comparison***

PhD Thesis

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A C A D E M I C Y E A R S 2 0 1 5 / 2 0 1 8

Abstract

Wheat, due to its adaptability to a wide range of environments and for the unique functional properties of its flour, represents the most widely grown, processed, and consumed cereal by humankind of temperate regions. Most of the modern wheat genotypes are derived from old wheats and have appreciable properties in terms of grain yield. Wheat is also the causing factors of many adverse reactions, such as celiac disease, allergies and non-celiac wheat sensitivity (NCWS) in susceptible people. This has led to an increasing interest for the old wheat genotypes which are generally considered better tolerated than the modern ones, but without any scientific evidence. The aim of the present work is the qualitative and quantitative comparison by a proteomic approach of the metabolic protein fractions extracted from the mature kernel of two old Sicilian durum wheat landraces (*Russello* and *Timilia reste bianche*), and *Simeto*, an improved durum wheat variety, widely spread in Italy and other Mediterranean countries, chosen as representative of the most widely commercial *cultivars*. The qualitative comparison of the protein composition revealed a remarkable similarity between old and modern *cultivar*. The quantitative evaluation of the identified proteins shows that some proteins are differentially expressed in old and modern varieties.

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LIST OF ABBREVIATIONS

CD	Celiac Disease
NCWS	Non-Celiac Wheat Sensitivity
cv	cultivar
IgE	Immunoglobulin E
CM	Chloroform/Methanol-soluble
LTP	Lipid Transfer Protein
Serpin	Serine Proteinase Inhibitor
S-rich	Sulfur-rich
S-poor	Sulfur-poor
HMW	High Molecular Weight
MALDI	Matrix-Assisted Laser Desorption/Ionization
TOF	Time Of Flight
RP-HPLC	Reverse Phase-High Performance Liquid Chromatography
nESI	nano ElectroSpray Ionization
EI	Electron Impact
Q	Quadrupole
IT	Ion Trap
CI	Chemical Ionization
CID	Collision Induced Dissociation
HCD	High-energy Collision Dissociation
ETD	Electron-transfer dissociation
LC	Liquid Chromatography
Q-OT-qIT	Orbitrap Fusion Tribrid
AUC	Area Under the Curve

MuDPIT	Multidimensional Protein Identification Technology
PSM	Peptide spectral matches
Pfam	Protein Family
FDR	False Discovery Rate
BLAST	Basic Local Alignment Search Tool
GO	Gene Ontology
PANTHER	Protein ANalysis THrough Evolutionary Relationship
DTT	dithiothreitol
IAA	iodoacetamide
FA	Formic Acid
ACN	Acetonitrile

1.Introduction

Wheat is one of the most important cereal for the humankind. In 2016, the global production of wheat exceeded 749 million tonnes, making it the second most-cropped cereal after maize.¹ Wheat contributes between 20% and 50% of the total calories in countries where this crop is cultivated, USA, China, Russia, India, Pakistan, Canada, Argentina, Australia and some countries of European Union, but the consumption of wheat is also increasing in countries where there aren't specific climate conditions, such as Sub-Saharan Africa, and particularly in countries undergoing urbanization.² Wheat is not only a source of calories but it also contains essential amino acids, minerals, vitamins and bioactive compounds for the human diet.² The importance of wheat depends on the possibility to transform its seeds into flour, semolina, etc., which give origin to a lot of food products and make substantial contributions to the dietary intake of energy and consequent impact on human health.³ All types of wheat are included in the grass *Poaceae* family and in particularly they belong to the genus *Triticum*.³ The most common varieties of this crop are *Triticum aestivum* ("common", "bread" or "soft" wheat) and *T. durum* (durum wheat).¹ The first one is an hexaploid species (AABBDD) whereas the second one is tetraploid with this genome composition AABB (Fig. 1).

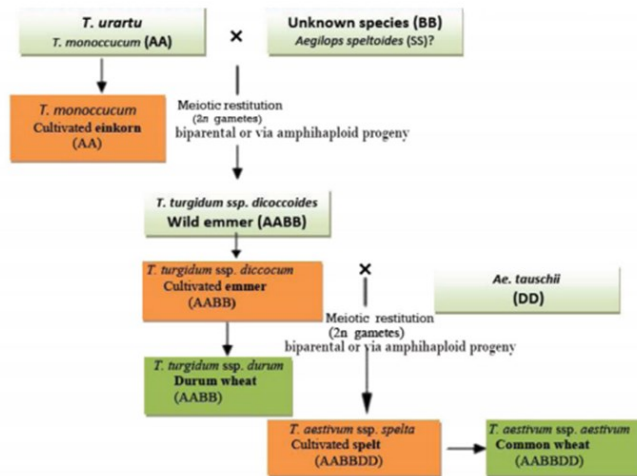


Figure 1. Phylogeny of domesticated species of the common wheat varieties⁴

In particularly the durum wheat is largely cultivated in the Mediterranean areas and predominantly used for making pasta. The main components of the wheat kernel are carbohydrates (70 – 80% of dehydrate flour), lipids (1.5 – 2.5% of dehydrate flour) and proteins (8 – 18% of dehydrate flour).

Wheat is also the causing factors of many adverse reactions, such as celiac disease (CD), allergies and non-celiac wheat sensitivity (NCWS) in susceptible people. At the present two main groups of wheat allergens are known: gliadin/glutenin (salt-insoluble proteins) and albumin/globulin (salt-soluble proteins) fractions. The first fraction is caused of celiac disease, a condition that results in a chronic inflammatory of the gastrointestinal tract, which affects the small intestine, causing atrophy of the absorbent apparatus and malabsorption of nutrients.⁵ Nowadays, the only treatment available for this condition is a lifelong diet gluten-free. On the other hand, albumins and globulins are cause of wheat allergies, that is the result of immunological reactions, mediated by allergen-specific immunoglobulin E (IgE). A typical example of wheat allergy is the Baker's asthma, a typical condition in which water soluble flour proteins bond to serum IgE as a result of inhalation of flour particles.³ In particularly, the most important allergens

are the α -amylase/trypsin inhibitor subunits, a class of hydrolytic enzymes, which catalyze the cleavage of the α -1,4 glycosidic linkage, and are widely diffused in nature. The plant α -amylase inhibitors exist in different polypeptides such as monomers of about 12 kDa, homodimers of 24 kDa and heterotetramers of about 60 kDa. The wheat tetrameric inhibitors are constituted by three different subunits belonging to the class of the CM (Chloroform/Methanol-soluble) proteins, which typically result in 13–15 kDa polypeptides under dissociating conditions.⁶ Recently, other several proteins have been linked to wheat allergy; in particular, lipid transfer protein (LTP), peroxidase, thioredoxin, serine proteinase inhibitor (serpin), thaumatin-like protein, acyl-CoA oxidase, fructosebisphosphate aldolase, triosephosphate isomerase, and glyceraldehyde-3-phosphate dehydrogenase.

In conclusion, NCWS shows similar symptoms to CD and wheat allergy, but without the serologic or histologic evidence of CD.⁷ It has also been defined as a variety of immunological, morphological, or symptomatic manifestations that is possible to observe after the ingestion of gluten in individuals in whom CD has been excluded.⁷ Nowadays, there isn't a specific test to reveal the diagnosis of NCWS; the diagnosis of this condition is most the time achieved by elimination: it is not CD and it is not wheat allergy.⁸ There are two important differences from people suffering of CD and people with NCWS. NCWS patients have the possibility to reintroduce the gluten in their diet and do not appear to be at risk for nutritional deficiencies, whereas CD patients have to follow a lifelong gluten-free diet.⁷

Beside to people who need to follow a gluten-free diet for health reasons, a new segment of consumers who consume gluten-free products as a lifestyle choice has arisen. Currently, a consistent percentage of the general population considers oneself to be suffering from problems caused by wheat and/or gluten ingestion, even if they are not celiac. This has led to an increasing interest for the old wheat genotypes, but the nutritional comparison between ancient and modern wheat varieties is still controversial. In literature, it is reported that the health benefit of ancient grains is not related to a single compound but to a general composition of these varieties.¹ In particular, ancient wheat species suggest that they could present a healthier and a better nutritional profile than modern wheats, because rich in vitamins, minerals and nutraceutical compounds.¹ The wheat cultivation was the first step in the transition from hunting of food to settled

agriculture, about 10,000 years ago. Einkorn and emmer were the earliest cultivated forms, which are diploid (AA) and tetraploid (AABB) species, respectively. Modern durum wheat was developed from the same wild ancestor as emmer. Now, emmer and durum are regarded as forms of the same species (*Triticum turgidum*). *Triticum aestivum* (bread wheat) goes back about 9,000 years ago by hybridization of cultivated emmer with wild “goat grass” (*Triticum tauschii*). Today, bread and durum wheat are the varieties more cultivated, but some “ancient wheats” (emmer, einkorn, etc.) continue to be produced in small amounts to satisfy the increasing demand for the health food market.² Although there is no precise definition, it is generally accepted that ancient wheat has remained unchanged over the last hundred years. On the contrary the modern wheats were modified and subjected to the “Green-Revolution”. This revolution was a mix of research and technological transfer initiatives that occurred between the 1930s and the late 1960s. The Green- Revolution was initiated by Strampelli, who was among the first, in Europe and in the World, to systematically apply Mendel's laws to modify some proprieties of this crop. At the end of this revolution modern varieties characterized by higher yield, a reduced susceptibility to diseases and insects, an increased tolerance to environmental stresses, a homogeneous maturation (to optimize harvest) were obtained. By using these modern varieties it was possible to increase the production of wheat with a concomitant decrease in genetic variability as well as a gradual impoverishment of the nutritional and nutraceutical properties.¹ On the contrary, the ancient varieties are cultivated with environmentally sustainable organic agriculture. The most common ancient wheat species commercially available are einkorn (*Triticum monococcum*), emmer (*Triticum dicoccum*), khorasan (*Triticum turgidum ssp. turanicum*) and spelt (*Triticum spelta*). In addition, there are several old cultivars of both *Triticum aestivum* and *Triticum durum* that remained unchanged over the years, namely *Russello*, *Senatore Cappelli*, *Timilia or Tumminia* and *Urria* (*Triticum durum*), as well as *Autonomia B*, *Frassineto*, *Gentil Rosso*, *Inallettibile*, *Maiorca*, *Sieve*, *Solina*, and *Verna* (*Triticum aestivum*).¹ The characteristic nature, low-input and organic managements, have increase the interest on ancient wheat cultivars, and for these reasons they are considered more “natural” with respect to modern wheats. Today, research performed on ancient wheat varieties is scarce and in particular

exhaustive proteomic comparisons of these species with respect to the modern wheat varieties are lacking.

1.1 Classification of wheat proteins

The classification of wheat proteins is based on their solubility and extractability in various solvent, named Osborne's classification (Fig. 2). By using this Osborne's classification, these proteins are grouped in:

- albumins, which are soluble in water;
- globulins, which are soluble in dilute salt solution;
- gliadins, which are soluble in 70% ethyl alcohol, and;
- glutenins, which are soluble in dilute acid solution or 70% ethyl alcohol under reducing conditions.³

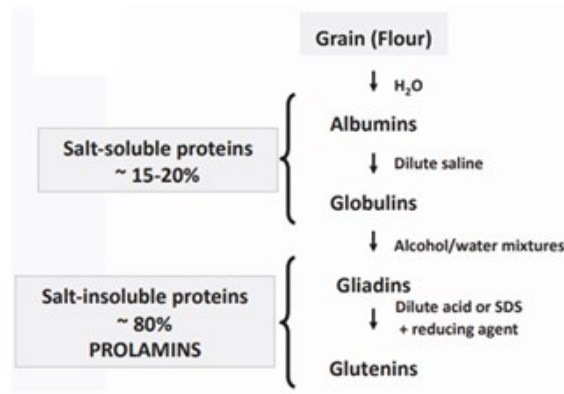


Figure 2. Protein fractions from wheat grain obtained by sequential extraction in different solvent⁹

The terms “albumins” and “globulins” are also more widely used for proteins with similar solubility properties from other organism while the prolamins are named specifically in different cereals: gliadins in wheat, hordeins in barley, secalins in rye, and zeins in maize.

Another classification of the cereal proteins is based on their functions: structural and metabolic proteins, protective proteins and storage proteins. The first group include a wide number of proteins, which contribute to cell, structure and function, such as enzyme, proteins involved in transport, growth, gene transcription and protein synthesis. This group include albumins and globulins of Osborne’s classification. The group of protective proteins comprises components involved in defence mechanism of the plant. The last group includes proteins deposited in seeds. These proteins provide a store of amino acids and carbon skeletons for germination and represent more than 50% of the total protein content in the mature cereal seeds.

Today the cereal proteins are classified into superfamilies: prolamins and cupins (Fig. 3). The prolamins superfamily includes seed proteins mainly, which can be divided into two types low-molecular-mass sulfur-rich seed proteins and the prolamins themselves. The proteins of the first group have molecular mass of about 7-16 KDa. This group include three major classes of cereal food allergens, 2S albumins, non-specific lipid transfer proteins and cereal α -amylase/trypsin inhibitors. The prolamins themselves shows a large variety of structure and properties. A classification, based to structure, divides these proteins in three groups: sulfur-rich (S-rich), sulfur-poor (S-poor) and High Molecular Weight (HMW) prolamins. The proteins of cupin superfamily are divided in germins and storage globulins. They are involved in the response to various stress conditions. The storage globulins, on the basis on their sedimentation coefficient, can be divided into 7S vicilin-type globulins and the 11S legumin-type globulins.¹⁰

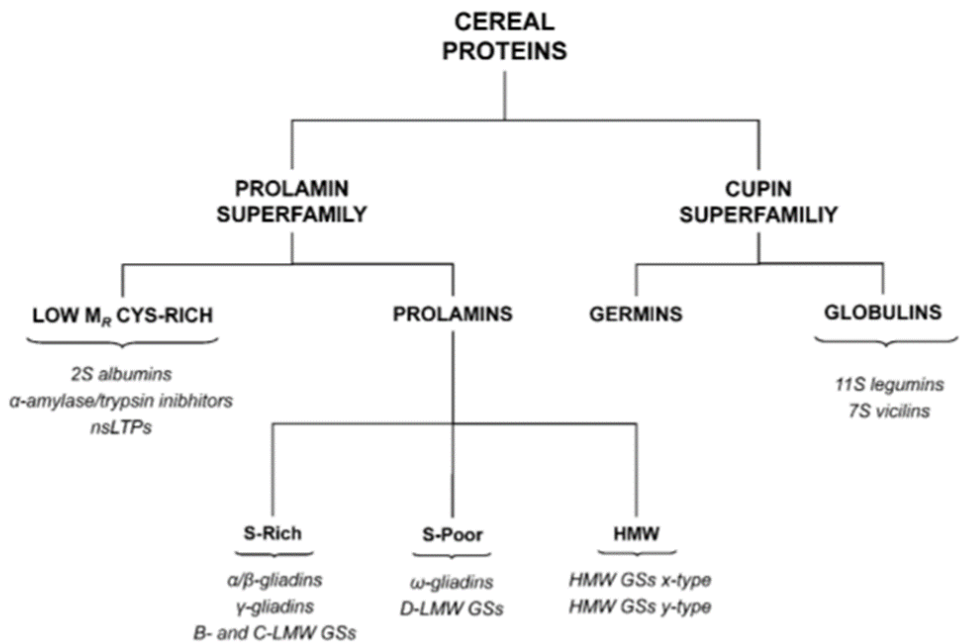


Figure 3. Classification of wheat proteins based to their structural and evolutionary relationships¹⁰

2. Proteomic analysis

The term "proteomics", proposed by analogy with the word genomics, indicates study in large-scale of proteome. Proteomics is more complex to genomics for various reasons. One reason is that several proteins may come from the expression of a single gene. Today, is almost never valid the old idea that to a gene corresponds a protein. Phenomena such as splicing and numerous post-translational modifications that characterize the mature product, may take place from the gene transcription to the "final protein product", which increase the variability of a single protein species.

There are three different proteomic approaches: "Bottom-up", "Shotgun" and "Top down" (Fig. 4) methods. In the first approach a complex mixture of proteins is separated by mono or two-dimensional

electrophoresis. Subsequently, the protein of interest is excised from the gel and subjected to enzymatic digestion. This is followed by mass spectrometry analysis (MALDI-TOF or RP-HPLC/nESIMSMS) and bioinformatics research. The second approach is commonly named "Shot-gun" or "MuDPIT" (Multidimensional Protein Identification Technology). By using this method, the mixture of proteins is directly subjected to enzymatic digestion and peptides are separated by on-line chromatography coupled to tandem mass spectrometry.¹¹ In this case, the experimental procedure is more easy but the processing of the data is difficult for the complexity of the system. The "top-down" approach consists in the study of intact protein ions and their direct fragmentation within the instrument without previous digestion¹².

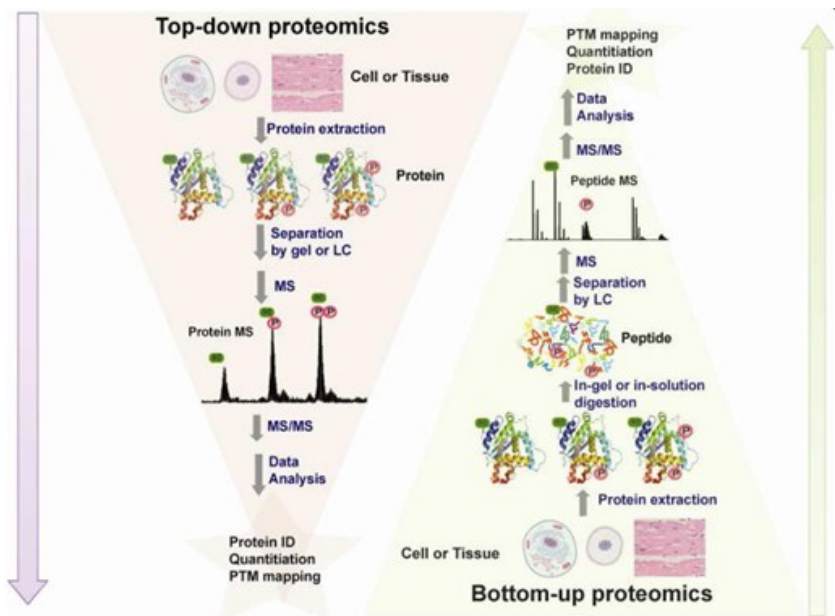


Figure 4. Schematic illustration of the difference between “top-down” and “bottom-up” proteomics

2.2 High Performance Liquid Chromatography (HPLC)

High-Performance Liquid Chromatography (HPLC), allows to perform separations of very complex mixtures. The principle of separation of this

technique is based on the formation of a pseudo-equilibrium between each component of the sample, the stationary phase and the liquid mobile phase flowing between the particles of the stationary phase. The dimensions of the particles, which thus confer a high efficiency, of the stationary phase are comprised between 3 and 10 μm . At the head of the column there is a pump to apply a pressure to elute the liquid mobile phase. In this way, the separation is faster, but the process takes place through a larger number of theoretical plates, which results in better resolution. Usually, the reverse-phase chromatography (RP-HPLC) is the common type of liquid chromatography used for the separation of protein and peptide mixtures. This technique uses apolar stationary phase, consists of more or less long alkyl chains (C4, C12, C18), linked to small spheres of silica and polar mobile phase, generally a mixture of two or more different solvents, whose flow rate is regulated by the respective pump. This makes possible to work in two different conditions of flow:

- isocratic conditions, where there is the same mobile phase composition during the analysis;
- conditions of gradient elution; in this case the solvent composition and consequently polarity are variable during the analysis. In this way, the gradient separates the analytes contained in the mixture in function of their affinity for the specific mobile phase compared to the stationary phase.

When a separation is effected with elution gradient, at the beginning of the separation the mobile phase is rich in the more polar solvent and, thereafter, the solvent with non-polar characteristics is increased over the time. Usually, in RP-HPLC water is the polar solvent and methanol or acetonitrile are the apolar organic solvents. In this way, initially the more polar components of the mixture are eluted, whereas the more apolar ones, which have a greater affinity for the organic eluent are eluted later. The main components of a modern HPLC are (Fig. 5):

- containers for solvents with degassing system;
- pumps, used to apply high pressures to elute the mobile phase in the system. The pump allows to maintains a stable and reproducible flow during the analysis;

- system for the introduction of the sample (sampling loop);
- column, in which the separation of the analytes occurs; the HPLC columns can have different dimensions and characteristics depending on the type of analyses to be performed, on the system used, the type of detector and, not least, the amount of sample available;
- detector, which generates a signal when reached by the eluted components of the mixture. The selection of the detector depends on the needs dictated by the nature of the sample, because universal and highly sensitive detectors for HPLC do not exist. Usually, the detectors used for liquid chromatography are based on the measure of the absorption of ultraviolet or visible light by the sample. For instance, detection of proteins is carried out at 220-224 nm. On the other hand, a particularly sensitive and versatile detector is represented by a mass spectrometer with electrospray ionization, which today is widely used in proteomic studies.

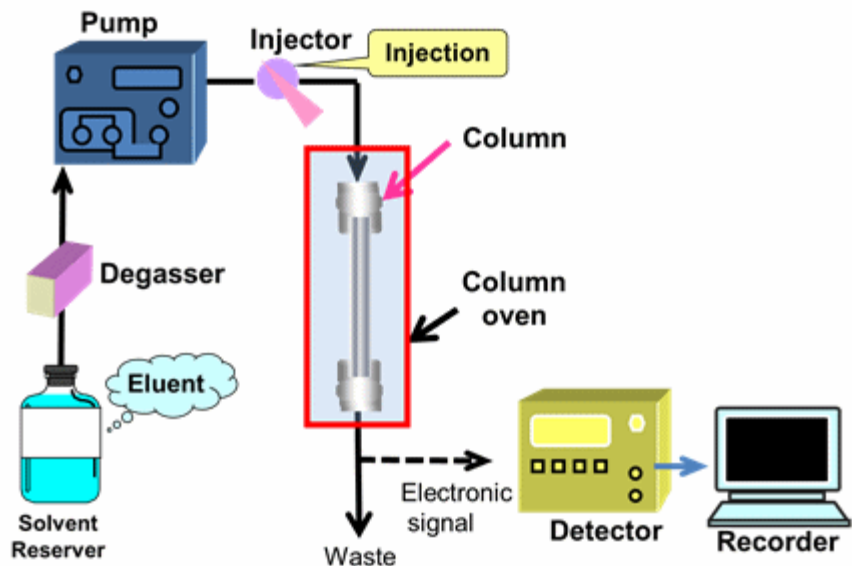


Figure 5. Main components of an HPLC system

2.3 Mass spectrometry

Mass spectrometry is an analytical technique based on the production of ionized molecules, the subsequent separation on the basis of their different mass/charge ratio (m/z) and detection of the ions produced. At the end, a graph of relative abundance versus m/z ratio is obtained. The principal constituents a mass spectrometer are (Fig. 6):

- system for the introduction of the sample;
- source, where the ionization of the sample occurs;
- analyzer, which performs a separation of the ions produced in the source according to their m/z ratio;
- detector, where the separated ions are detected;
- vacuum system, whose task is to keep the various parts of the instrument under vacuum, the presence of which (the pressure is around 10^{-6} - 10^{-8} Torr) is needed primarily to avoid the collision of the ions produced with the atmospheric gases.

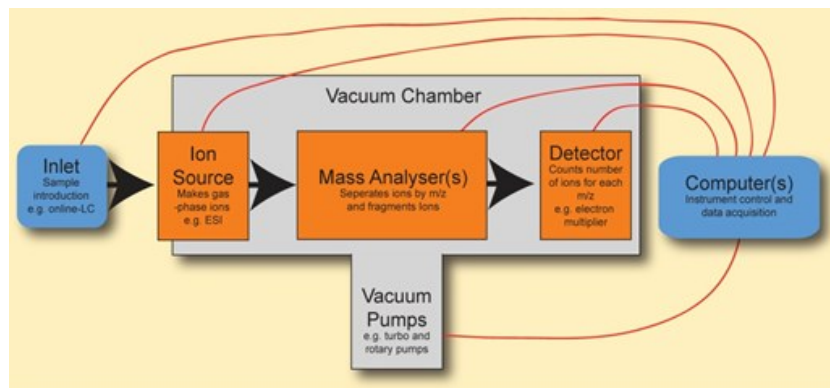


Figure 6. Block diagram of a mass spectrometer

2.3.1 Ion sources

The electron impact (EI) and the chemical ionization (CI) were the first sources for ionization of the sample, suitable only for molecules with low molecular weight and easy to transfer in the gas phase. In the late 1980, thanks to the introduction of two new methods for desorption/ionization, respectively known as MALDI (Matrix-Assisted Laser Desorption/Ionization) and ESI (Electrospray Ionization), mass spectrometry has assumed an important role also in the study and characterization of biomolecules.

In particular, the electrospray ionization (ESI) is a soft ionization technique since it does not produce fragmentation of the sample. This ionization technique is the ideal interface for the on line coupling of a chromatographic system (RP-HPLC/ESI MS) and a mass spectrometer, and assumed an important role in the field of mass spectrometry for the ability to bring into gas phase and ionize macromolecules of biological origin. Electrospray mass spectrometry (ESI-MS) allows to obtain, from a solution of analyte introduced into the source by direct infusion or coming from a chromatographic column, single-charged ions and multiple charged ions which are thus sent to the analyzer and to the detection system.

By using a capillary tube of silica the protein solution is introduced into the source. Inside the ionization chamber, a spray is produced between the metallic tip of the needle and a counter electrode, where it is present a strong electric field (3-5 kV) that disperses the solution emerging from the needle into an aerosol of droplets with a high charge concentration. The desolvation of the droplets of the spray is obtained by using a stream of nitrogen suitably heated or just the high temperature of the capillary tube. The generally used solvent is water mixed with an organic solvent (acetonitrile, methanol, or propanol) and small amounts of or a weak acid (trifluoroacetic acid, acetic acid or formic acid) or a weak base (ammonia solution) to facilitate the ionization of the sample and the formation, respectively, of positive or negative ions. The mechanism through which the ions are formed starting from the charged drops of sample has not yet been completely

clarified; several models have been proposed, including a qualitative model compatible with the mechanisms proposed by Smith, Fenn and Röllgen.^{13,14,15} According to this model, in a first moment the formation of micro-droplets fillers whose dimensions are related to their surface tension is observed; the hot gas stream causes the desolvation of these micro-droplets, tending to bring together the charged molecules. When the force of the Coulomb repulsion equals the surface tension of the droplet (Rayleigh limit), it explodes producing other smaller droplets (nano-droplets)¹⁶ that are subject to further desolvation (Fig. 7).

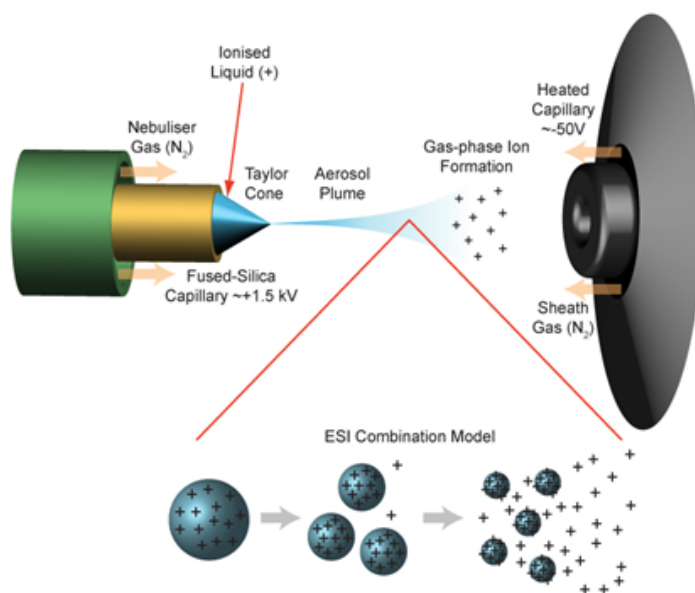


Figure 7. ESI source and model of ions formation

The pre-chamber is located at a pressure of 10^{-1} - 10^{-2} Torr, only a part of the ions arrives to this part of the instrument. Subsequently, the ion beam is focused, through a series of electrostatic lenses (skimmers), and reaches the analyzer (10^{-6} - 10^{-7} Torr), where separation takes place based on the value of the m/z ratio. The formation of multiple charged ions allows to display ions with high masses even working with analyzers that have limited mass range and, therefore, makes this ionization method an excellent tool for the analysis of peptides and proteins. A

typical ESI spectrum of positive ions of a protein consists of a set of peaks, each of which is generated from the analyte that has linked a specific number of protons. The proteins are usually analyzed as positive ions because a series of multi-charged protein ions generated in the source is mainly related to the protonation of basic sites of molecules. In general, in a protein, the number of basic amino acid residues determines the maximum number of protons that the molecule can take. The ESI spectrum of small molecules shows a precise correlation between the number of basic sites present in the structure and multi-charged ions. When the size of the molecule increases, this correlation is not so rigorous because some of the basic sites will be located inside the protein itself according to a particular conformation, and will be protonable with difficulty. The capacity to protonate a protein of high molecular weight is closely related to the conformation that the protein assumes in solution under the experimental conditions (pH, temperature, presence of denaturing agents). ESI mass spectrometry constitutes a particularly powerful and versatile detector for high performance liquid chromatography (HPLC). Tandem mass spectrometry is employed in order to select an ion with a given m/z ratio ("precursor" ion) and subsequently to fragment it; fragmentation leads to the formation of lower mass ions ("fragment" ions), which are analyzed in a second stage of analysis of mass ¹⁷. The characteristic fragmentation peaks in the MS/MS spectra allow to obtain important information on the molecular structure of the precursor ion. In the case of peptides, the fragment ions are generated by cleavage of the peptide bond with retention of the positive charge at the N-terminal (b series) or in the C-terminal part (γ series) along the main chain (Fig. 8), and allow to go back to the amino acid sequence of the precursor peptide.

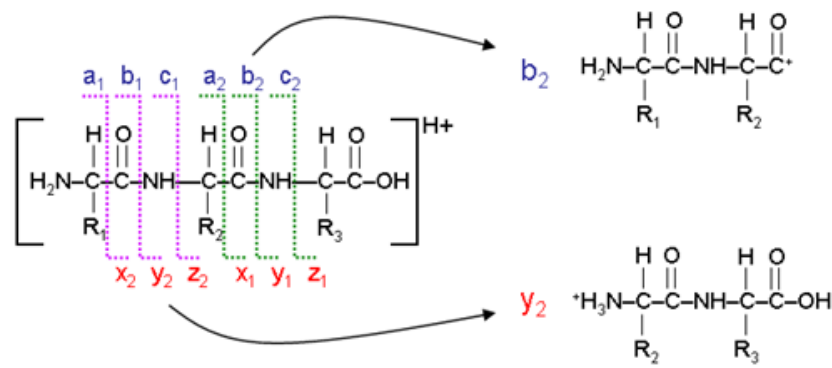


Figure 8. Scheme of the typical peptide fragmentation

2.3.2 Mass analyzer

The ion source can be interfaced with different mass analyzers. The most commonly used are quadrupole (Q), ion trap (IT), time-of-flight (TOF) and Orbitrap. The characteristics of these mass analyzer are different both in principles of operation and performance.

In particular, in 1999 Alexander Makarov developed the “Orbitrap” (Fig. 9), a new mass analyzer constitutes by an inner electrode (central) and external electrode, axially symmetrical, which create a combined square logarithmic electrostatic potential.

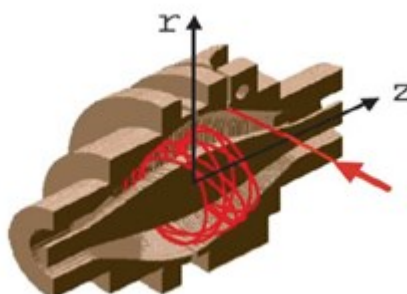


Figure 9. Ion motion within an Orbitrap analyzer

The ions rotate around a center electrode and oscillate with harmonic motion along its axis (z direction) with a frequency characteristic of their m/z values. As mentioned, within this analyzer,

the axial symmetric electrodes create a square logarithmic U (electrostatic potential), which can be calculated through the equation:

$$U(r,z) = \frac{k}{2} \left(z^2 - \frac{r^2}{2} \right) + \frac{k}{2} (R_m)^2 \ln \left[\frac{r}{R_m} \right] + C$$

where r and z are the cyclic coordinates, C is a constant, k is the field curvature and R_m is the characteristic radius. In this U field, a rotational motion around the electrode and an oscillatory motion along the axes create stable trajectories of the ions, which result in a complex spiral. The equations that describes this motion for this mass analyzer are very complex. From these equations it follows that the mass and the charge are correlated with the frequency of axial oscillations, expressed in radiant/second:

$$\omega = \sqrt{(q/m)k}$$

ω is completely independent of the energy and position of the ions, and thus can be used for analysis of mass (in fact in the expression appears the ratio q/m). All ions have then a harmonic oscillatory motion of the same amplitude but of different ω frequency. These frequencies are measured in a non-destructive way by a differential amplifier, which acquires the signals of the current image in the time domain. For each ion is produced a wave function; therefore, a mixture of ions gives rise to overlapped signals that can be converted to a mass spectrum thanks to Fourier transform.

In 2013, a new instrument was introduced, the Orbitrap Fusion Tribrid Mass Spectrometer (Fig. 10). This instrument combines the best of quadrupole, linear ion trap and Orbitrap mass analysis in a new instrument. The resolution of this instrument is up to 450,000 FWHM. Moreover, the precursor selection using a quadrupole mass filter allows the ion trap and Orbitrap mass analyzer to operate in parallel for

excellent sensitivity and selectivity. Also, multiple dissociation techniques (CID, HCD and ETD) are possible.

Collision Induced Dissociation (CID) is the most commonly method of fragmentation in proteomics.¹⁸ By using anelastic collision, selected precursor ions are collided with an inert gas. CID fragmentation occurs at the peptide bond between the carboxyl group and amino group. The produced fragments are γ -ions and b-ions.¹⁹ High-energy Collision Dissociation (HCD) is a fragmentation method which produces the same γ /b-ions and Y/B-ions as CID. It can be performed only in instrument with HCD fragmentation cell and uses higher energy than CID. The theory of precursor ion fragmentation the in Electron-transfer dissociation (ETD) is still debated, but it is known that ETD produces fragments of c/z-type, given complementary information about peptide sequence.

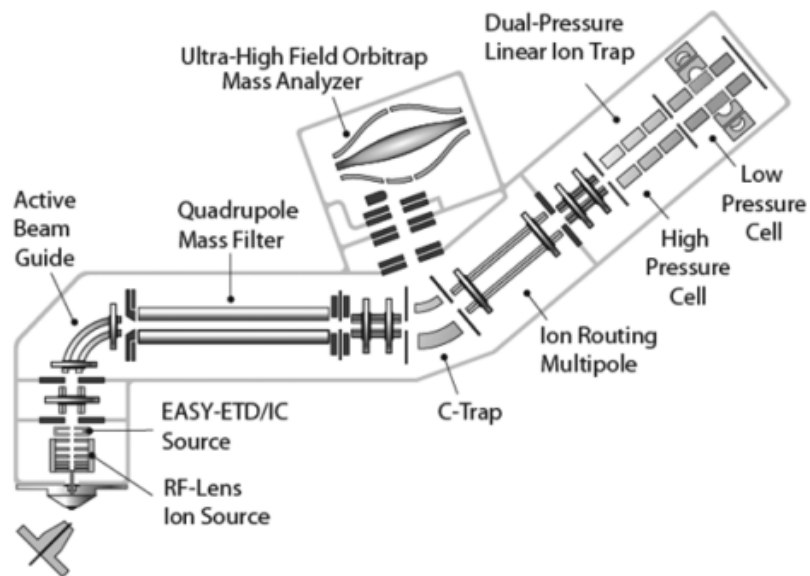


Figure 10. Schematic representation of the Fusion instrument²⁰

2.4 Bioinformatic search

Adequate support of software to analyze the collected data is fundamental in proteomic analysis. Today, there are a variety of algorithms for the interpretation of peptide fragmentation data. LC-MS/MS data in this work were processed using PEAKS 8.5 (Bioinformatics Solutions Inc.) software.

2.4.1 Peaks 8.5 (Bioinformatics Solutions Inc.)

In order to identify proteins, LC-MS/MS data were processed using PEAKS 8.5 (Bioinformatics Solutions Inc.). Usually, the algorithm to identify the identity of a peptide compare the fragmentation pattern to theoretical fragmentation pattern derived from protein sequence databases. PEAKS Q 8.5 (Bioinformatics Solutions Inc.) employed in the present work to process LC-MS/MS data for proteins identification uses this method and integrates database search with *de novo* sequencing for peptide identification. *De novo* sequencing derives directly the peptide sequence from the MS/MS spectrum without the need of a sequence database.²¹

Label-free quantification is included in the PEAKS Q module. It is used in the study of large scale proteomics to obtain a fast protein profiling. This quantification method is based on the detection of peptide features (mass, retention time and signal intensity) in multiple samples. For each sample is obtained a feature detections and than by using the EM (expectation-maximization) algorithm, these features can be overlapped. The features of the same peptide from different samples are aligned together using a high performance retention time alignment algorithm.²²

3. Label-free proteomic analysis

Mass spectrometry plays an important role in proteomic analysis. The new techniques, developed in recent years, gel-free based “shotgun” proteomic, such as Multidimensional Protein Identification (MudPIT) allow to study the protein expression in complex biological system.^{23,24} Proteomic studies can be performed to obtain both absolute (using internal standards) or relative quantification by different techniques including label-based and label-free approaches.

In label-based approach a stable isotope is used to label the sample by biosynthetic or chemical reactions.²⁵ Labelling strategies are often preferred because they are considered more accurate in quantitating protein abundances. However, this technique requires expensive isotope labels, specific software and expertise to analyse data.³² Moreover, most of the label-based methods require more steps in sample preparation and higher sample concentration, are more expensive and can only be performed for a limited number of samples.^{26,27}

MS-based label-free quantitative proteomics avoids the use of isotopes to label the samples under investigation and this approach can be used in “shotgun” analysis (analysis of the whole proteome) or in targeted analysis (analysis of specific proteins) and it can be applied when labelling is not possible.²⁸ There is a correlation between protein abundance and peaks areas^{29,30} or number of MS/MS spectra.³¹ Today, label-free methods are divided in two groups: (i) measurement of the intensity of the ion precursor signal or area under the curve (AUC) and (ii) spectral counting, which is based on counting of the number of peptides assigned to a protein in an MS/MS experiment (Fig.

11).³²

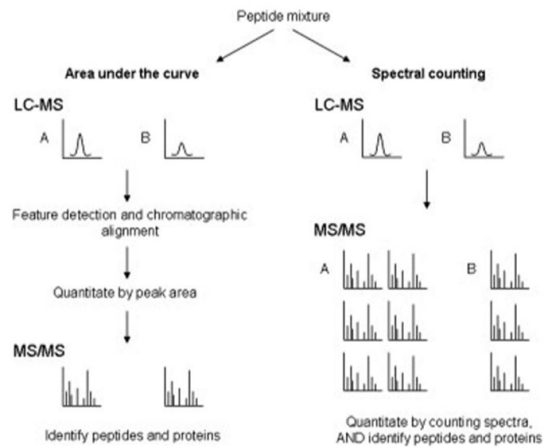


Figure 11. Schematic diagram of the two main approaches used in label-free proteomics. AUC quantitation is performed at the MS level and peptides found to have differential expression are identified by MS/MS. In Spectral counting quantitation and identification are performed simultaneously at the MS/MS level.³²

Regardless of which label-free quantitative proteomics method is used, the analysis includes the following fundamental steps:

- sample preparation including extraction, reduction, alkylation and digestion;
- sample separation using liquid chromatography and ESI-MS/MS;
- data analysis, including protein identification, quantification and statistical analysis.

After the acquisition of MS/MS spectra, the raw data need to be processed by a software. Label-free proteomics software workflows typically consist of multiple steps: peptide peak picking, peptide identification, feature finding, matching of the features with identified peptide, alignment of the features in different samples. Protein quantifications is finally obtained from quantified peptides.^{33,34}

4. Aim of work

Wheat flour is the main ingredient in most type of breads, pastries and pastas worldwide, because of its unique protein composition.² However, as above reported, wheat is also the causing factors of many adverse reactions, such as celiac disease, allergies and non-celiac wheat sensitivity in susceptible people. A consistent percentage of the general population considers oneself to be suffering from problems caused by wheat and/or gluten ingestion, even if they are not celiac. These people follow a gluten-free diet for health reasons as a lifestyle choice. Moreover, the close relationship between diet and health is now generally recognized and the increased wellness and consciousness, especially in developed Countries, has led to an increasing interest for the old wheat genotypes, which are generally considered better tolerated than the modern ones, but without any scientific evidence.

This work is aimed to the comparative proteomic analysis and a quantitative evaluation by using a MS label-free approach of the metabolic proteins in old ad modern durum wheat varieties. Proteins were extracted from the mature kernel of *Russello* and *Timilia reste bianche*, two old Sicilian durum wheat landraces, and *Simeto*, an improved durum wheat variety, widely spread in Italy and other European countries (Spain, Greece, etc.). *Russello* and *Timilia* are cultivated in areas such as Agrigento, Caltanissetta, Ragusa e Trapani. They are used in the production of high digestible bread and pasta. *Simeto* is a durum wheat variety, common in southern Italy and in islands, which has an excellent capacity to adapt to different temperature conditions. The protein profile of this variety can be considered representative of the commercial cultivars most widely used in the current practice.

To improve the qualitative comparison between these *cultivar* an enrichment of two fractions, metabolic and CM-like proteins, was carried out. Although the separation of these protein fractions is not selective and consequently a cross contamination between the two fractions is observed, this method allows to enhance the number of protein identifications in each fractions.

The quantitative evaluation was performed by using a “shotgun” approach on the whole metabolic protein fraction, without any separation. It was not possible to use the same extracts to carry out the qualitative and quantitative comparison because the cross contamination undermined the quantitative results.

5. Material and methods

5.1 Materials

The wheat flour of *Russello*, *Timilia Reste Bianche* and *Simeto* were provide from Cereal Research Centre (CREA), Foggia.

All chemicals were of the highest purity commercially available and were used without further purification. KCl, NaCl, K₂HPO₄, acetone, methanol and Tris-HCl were purchased from Carlo Erba (Milan, Italy). Formic Acid (FA), Protease Inhibitor Cocktail, EDTA, ammonium acetate, dithiothreitol (DTT), iodoacetamide (IAA) and lysozyme were obtained from Aldrich (St. Louis, Missouri, USA). Modified porcine trypsin was purchased from Promega (Madison, WI, USA). Water and acetonitrile (OPTIMA® LC/MS grade) for LC/MS analyses were purchased from Fisher Scientific (Milan, Italy).

5.2 Extraction of the metabolic and CM-like protein fractions for the qualitative comparison

Wheat flours (200 mg) were suspended in 2 mL of cold (4°C) extraction solution (50 mM Tris-HCl, 100 mM KCl, 5 mM EDTA, Protease Inhibitor Cocktail, pH 7.8) in order to extract the metabolic and CM-like proteins. The solution was incubated on ice for 5 minutes with intermittent mixing and centrifuged at 12000 rpm for 15 minutes at 4°C. The obtained soluble fraction was collected and five volumes of 0.1 M ammonium acetate in methanol was added. Following incubation over

night at -20°C, the solution was centrifuged at 3000 rpm for 15 minutes at room temperature. The pellet (constituted by the metabolic proteins) was collected and rinsed in 3 mL of ammonium bicarbonate buffer 0.1 M, pH 8.2. The proteins in the supernatant (CM-like proteins) were precipitated by addition of four volumes of cold acetone. The resulting mixture was kept overnight at -20°C and subsequently centrifuged at 3000 rpm for 15 minutes at room temperature. Finally, the pellet (CM-like proteins) was rinsed in 1.5 mL of ammonium bicarbonate buffer 0.1 M, pH 8.2.³⁵ The concentration for each extract was determined by fluorimetric assay.

5.3 In-solution digestion of metabolic and CM-like proteins

40 µL (about 60 µg) of each protein extract were reduced by DTT (3 hours, 20°C), alkylated with IAA (1 hour, 20°C) and digested by porcine trypsin overnight at 37°C.

5.3 Extraction of the metabolic proteins for the quantitative comparison

200 mg of wheat flour were extracted in 2 mL of extraction buffer (0.4 M NaCl, 0.067 M K₂HPO₄, pH 7.6) for 15 min under continuous stirring at 20°C. The insoluble fraction was spinned down at 12,000 x g for 15 min in an Eppendorf centrifuge. The pellet material was separated and the extraction procedure was repeated twice. The supernatants from these extractions were pulled and stored at -80°C until required.

5.4 In-solution digestion of the metabolic proteins

Lysozyme (0.8 µg) was added internal standard to 50 µg (about 50 µL) of each protein extract. Each sample was reduced by DTT (3 hours, 20°C), alkylated with IAA (1 hour, 20°C) and digested by porcine trypsin

overnight at 37°C. In each sample a 5% aqueous solution of formic acid was added to obtain a final volume of 2 mL.

5.5 Liquid chromatography and tandem mass spectrometry (LC-MS/MS) analysis

Mass spectrometry data were acquired on a Thermo Fisher Scientific Orbitrap Fusion Tribrid (Q-OT-qIT) mass spectrometer (Thermo Fisher Scientific, Bremen, Germany). Liquid chromatography was carried out using a Thermo Scientific Dionex UltiMate 3000 RSLCnano system (Sunnyvale, CA). One microliter of the reconstituted sample was loaded onto an Acclaim[®] Nano Trap C18 Column (100 μm i. d. x 2 cm, 5 μm particle size, 100 \AA). After washing the trapping column with solvent A (H_2O + 0.1% FA) for 3 min at a flow rate of 7 $\mu\text{L}/\text{min}$, the peptides were eluted from the trapping column onto a PepMap[®] RSLC C18 EASY-Spray column (75 μm i. d. x 50 cm, 3 μm particle size, 100 \AA). Peptides were separated by elution at a flow rate of 0.25 $\mu\text{L}/\text{min}$ at 40°C with a linear gradient of solvent B (ACN + 0.1% FA) in A, 5% for 3 min, followed by 5% to 20% in 32 min, 20% to 40% in 30 min, 40% to 60% in 20 min and 60% to 98% in 15 min. We finished by holding 98% B for 5 min, 98% to 5% in one minute and re-equilibrating the column at 5% B for 20 min. The eluting peptide cations were converted to gas-phase ions by electrospray ionization using a source voltage of 1.75 kV and introduced into the mass spectrometer through a heated ion transfer tube (275 °C). Survey scans of peptide precursors from 200 to 1600 m/z were performed at 120K resolution (@ 200 m/z). Tandem MS was performed by isolation at 1.6 Th with the quadrupole, HCD fragmentation with normalized collision energy of 35, and rapid scan MS analysis in the ion trap. Only those precursors with charge state 2–4 and an intensity above the threshold of $5 \cdot 10^3$ were sampled for MS. The dynamic exclusion duration was set to 60 s with a 10 ppm tolerance around the selected precursor and its isotopes. Monoisotopic precursor selection was turned on. The instrument was run in top speed mode with 3 s cycles, meaning that the instrument would continuously perform MS² events until the list of non-excluded precursors diminishes to zero or 3 s, whichever is

shorter. MS/MS spectral quality was enhanced enabling the parallelizable time option (i.e. by using all parallelizable time during full scan detection for MS/MS precursor injection and detection). Mass spectrometer calibration was performed by using the Pierce® LTQ Velos ESI Positive Ion Calibration Solution (Thermo Fisher Scientific). MS data acquisition was carried out by utilizing the *Xcalibur* v. 3.0.63 software (Thermo Fisher Scientific).

5.6 Database search

LC–MS/MS data were processed using PEAKS de novo sequencing software (v. 8.5, Bioinformatics Solutions Inc., Waterloo, ON Canada). Data were searched against a dedicated protein database (7612 protein sequences), including only entries of *Triticum*, *Oryza*, *Hordeum*, *Avena*, *Secale*, *Maize* and *Brachypodium* species from UniProt database (release July 2018).

Database search in the qualitative analysis was carried out using the following parameters: i) full tryptic peptides with a maximum of 3 missed cleavage sites; ii) cysteine carbamidomethylation as fixed modification; iii) oxidation of methionine, transformation of N-terminal glutamine and N-terminal glutamic acid residue in the pyroglutamic acid form as variable modifications. The precursor mass tolerance threshold was 10 ppm and the max fragment mass error was set to 0.6 Da. Peptide spectral matches (PSM) were validated using a Target Decoy PSM Validator node based on q-values at a 0.1% FDR. A protein was considered identified if a minimum of two peptides were matched. Proteins containing the same peptides and that could not be differentiated based on MS/MS analysis alone were grouped to satisfy the principles of parsimony.

Finally, the identified unique gene products were classified, when available, by their Gene Ontology annotation (biological processes and molecular functions).

Label-free quantification analysis was performed processing LC–MS/MS data by PEAKS Q (v. 8.5, Bioinformatics Solutions Inc., Waterloo, ON Canada). This quantification method is based on the detection of

peptide features (mass, retention time and signal intensity) in multiple samples. For each sample is obtained a feature detections and than by using the EM (expectation-maximization) algorithm, these features can be overlapped. The features of the same peptide from different samples are aligned together using a high performance retention time alignment algorithm.²⁴ Proteins were considered as differentially expressed, for fold change ratios > 2 and significance > 20 .

6. Results and discussion

6.1 Qualitative comparison

Three biological replicates for each *cultivar* were analyzed. The extraction procedure of metabolic and CM-like proteins was carried out one time for each biological replicate of each *cultivar*. The extraction of these protein fractions is not selective, therefore a cross-contamination between these protein groups was obtained. Then, each extract was subjected to in-solution digestion followed by triplicate RP-nHPLC/nESI-MS/MS analysis and database search, in order to assess the reproducibility of the available MS data. Fig. 12 show a scheme of the adopted procedure for each *cultivar*.

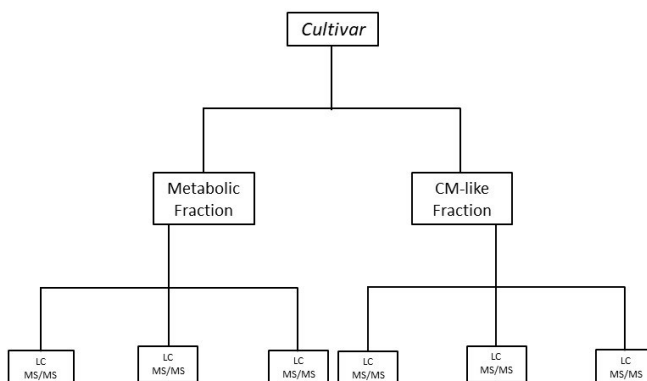


Figure 12. Scheme of the adopted analytical procedure for each *cultivar*

In order to obtain the two final lists of proteins (metabolic and CM-like proteins) for each variety here investigated, the following approach was adopted. Firstly, for each extract (i.e. metabolic or CM-like fraction) the lists of the proteins identified in the triplicate LC-MS/MS analyses were compared. Only those proteins identified at least twice were considered reliable. Then, the lists of proteins reliably identified in each biological replicate were compared, and only those proteins identified at least twice were considered to compile the final list for each *cultivar*.

The approach above described allowed the identification of 408 for *Russello*, 423 for *Timilia Reste Bianche* and 483 for *Simeto*, in their respective metabolic fractions. The lists of the identified proteins for each *cultivar* are reported in Tab. 1, 2 and 3, respectively. By the same approach, 100 proteins for *Russello*, 113 for *Timilia Reste Bianche* and 104 for *Simeto*, were identified in the CM-like fractions. The lists of the identified proteins for each *cultivar* are reported in Tab. 4, 5 and 6, respectively.

In order to perform a qualitative comparison among the *cultivar* investigated, the list of the identified proteins in the metabolic fraction of each ancient *cultivar* was compared with that obtained for the modern *cultivar* *Simeto*.

To carry out the Gene Ontology analysis for each *cultivar*, gene symbols were assigned to all the identified proteins by using the Uniprot Knowledge

database (<http://www.uniprot.org/>). It should be noted that the limited annotation of the wheat proteins in many cases required an additional step aimed to obtain coding gene information. So that, when the gene symbol was not available, the corresponding protein sequence was subjected to a sequence similarity search by BLAST (Basic Local Alignment Search Tool; <http://blast.ncbi.nlm.nih.gov/Blast.cgi>). Via this strategy, it was also possible to classify many “uncharacterized proteins” by finding homologous proteins present in databases, generally sharing more than 70% sequence similarity. By this approach, the proteins were grouped into protein families (unique gene products), taking into account, when available, the corresponding gene symbol and were subjected to gene ontology (GO) analysis through the PANTHER (Protein ANalysis THrough Evolutionary Relationship) system (<http://www.pantherdb.org>) by using the *Oryza sativa*, *Brachypodium* and *Arabidopsis thaliana* genome annotations as background.

A general qualitative comparison of the metabolic fractions of *Russello*, *Timilia* and *Simeto* revealed that 332 proteins are common to all *cultivar* (Fig. 13).

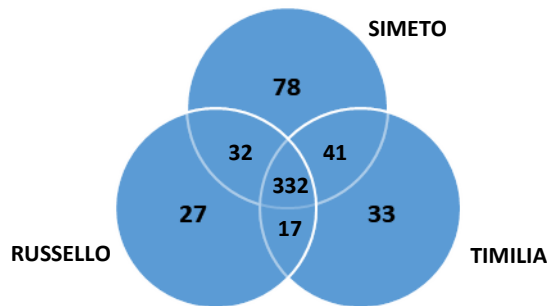


Figure 13. Venn diagram of the number of identified proteins in the metabolic fractions of *cultivar Russello*, *Timilia reste bianche* and *Simeto*

It was found that 78, 27 and 33 proteins are exclusive of the *cultivar Simeto*, *Russello* and *Timilia* respectively. The lists of these proteins are reported in Tab. 7, 8 and 9.

The identified metabolic proteins were classified in Molecular Function and Biological Process by Gene Ontology (Fig. 14). The lists of gene symbols used in the classification for each *cultivar*, *Russello*, *Timilia* and *Simeto*, are

reported in Tab. 10, 11 and 12, respectively. The qualitative proteomic analysis of these durum wheat varieties revealed a remarkable similarity in the protein composition between old and modern *cultivar*. In particular, this comparison reveals that most of them are involved in binding, structural molecule activity and catalytic activity and play a role in the same biological processes.

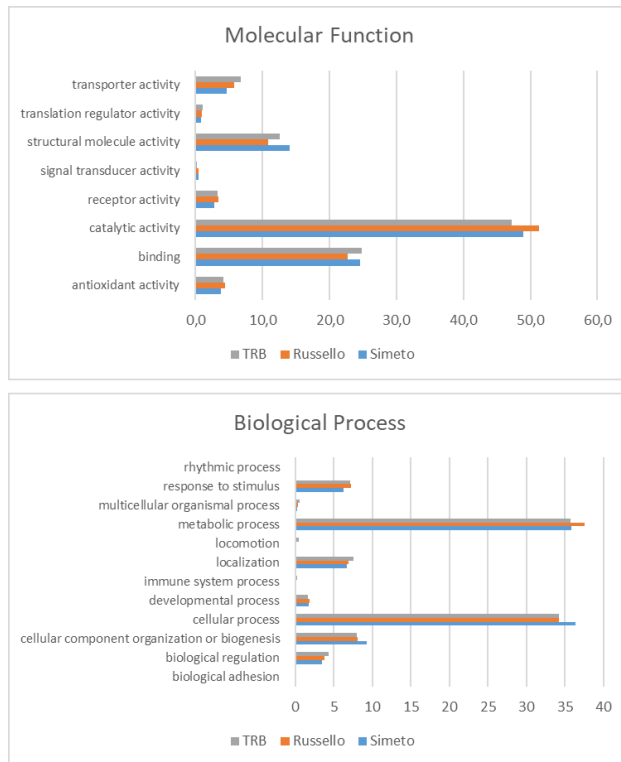


Figure 14. Gene Ontology of the metabolic proteins identified in *Russello*, *Timilia reste bianche* and *Simeto*

6.1.1 Qualitative comparison of the metabolic protein fractions of *Russello* and *Simeto*

Qualitative comparison of the metabolic protein fraction of *Russello* and *Simeto*, revealed that these two *cultivar* shared 364 proteins, whereas 44

and 119 are exclusively identified in the *cultivar Russello* and *Simeto*, respectively.

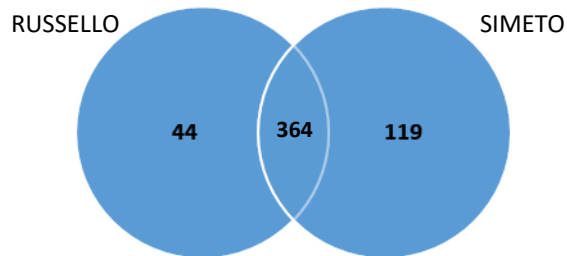


Figure 15. Venn diagram of the number of identified proteins in the metabolic fractions of *cultivar Russello* and *Simeto*

Gene Ontology analysis of the proteins identified exclusively in *Russello* and *Simeto*, allowed to ascertain their molecular function and the biological process in which they are involved (Fig. 16).

As depicted by the Molecular Function distribution, the proteins exclusively found in *Russello* or in *Simeto*, are mainly involved in catalytic activity, binding and structural molecular activity. The antioxidant activity is higher in the old variety (10.6%) respect to *Simeto* (3.8%). The proteins involved in this molecular function are different in the cultivars: *superoxide dismutase*, *peroxidase* in *Russello* and *L-ascorbate peroxidase* in *Simeto*, with only one protein, *catalase*, common to both varieties.

The comparison of Biological Process distribution does not show significant differences for almost all the biological processes reported, with the exception of the response to stimulus. Indeed, the *cv. Simeto* shows a percentage (5.5%) of proteins playing a role in this biological process which is lower with respect to that reported for the *cv. Russello* (14.3%). This biological process is divided in two sub processes: response to endogenous stimulus and response to stress. For the *cv Russello* there are two proteins involved in the first one (pyruvate kinase 1 cytosolic and ras-related protein RIC1) and seven proteins involved in the second one (belonging to the families *superoxide dismutase*, *peroxidase*, *catalase* and *heat shock protein*).

The proteins involved in the response to endogenous stimulus in *cv Simeto* is the *pyruvate kinase 2 cytosolic*, whereas the proteins involved in the response to stress are six proteins belonging to ribosomal proteins, L-ascorbate peroxidase, catalase and ubiquitin-activating enzyme.

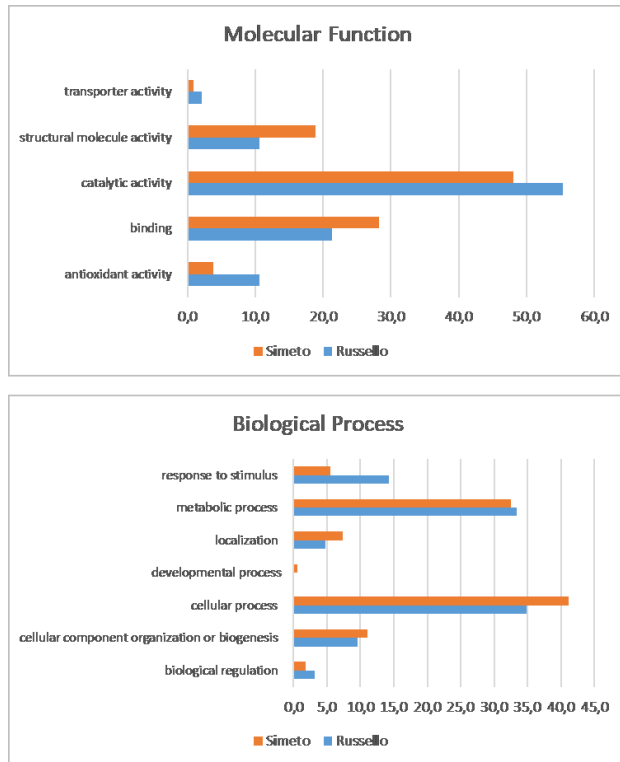


Figure 16. Gene Ontology of the unique proteins identified in the comparison *Russello* with *Simeto*

6.1.2 Qualitative comparison of the metabolic protein fractions of *Timilia reste bianche* and *Simeto*

The list of proteins identified in the metabolic fraction of *Timilia reste bianche* and *Simeto*, shared 373 proteins. On the contrary, 50 components were unique for the *cv Timilia* and 110 were exclusively found in the *cv Simeto*.



Figure 17. Venn diagram of the number of identified proteins in the metabolic fractions of *cultivar Timilia reste bianca* and *Simeto*

Gene Ontology analysis (e.g. classification based on their molecular function, biological process and protein class) of the unique proteins for these two cultivars are reported in Fig. 18.

Comparison of Molecular Function distributions, shows that most of the exclusive proteins of these two cultivars are mainly devoted to catalytic, binding and structural molecular activities. However, the *cultivar Simeto* also presents exclusively proteins involved in structural molecule activity (*tubulin alpha* and *beta*, *60s* and *39s ribosomal protein* family) and in receptor activity (*V-type protein* family), two molecular functions absent in exclusive proteins of *Timilia*. On the contrary, the *cultivar Timilia* is characterized for the presence of a higher antioxidant activity (12.2%) respect to 2% in *Simeto* but the proteins involved in this activity belong to the same families.

As above reported for the comparison between *Simeto* and *Russello*, the Biological Process distribution of unique proteins of *Timilia* and *Simeto* does not show significant differences for almost all the biological processes reported.

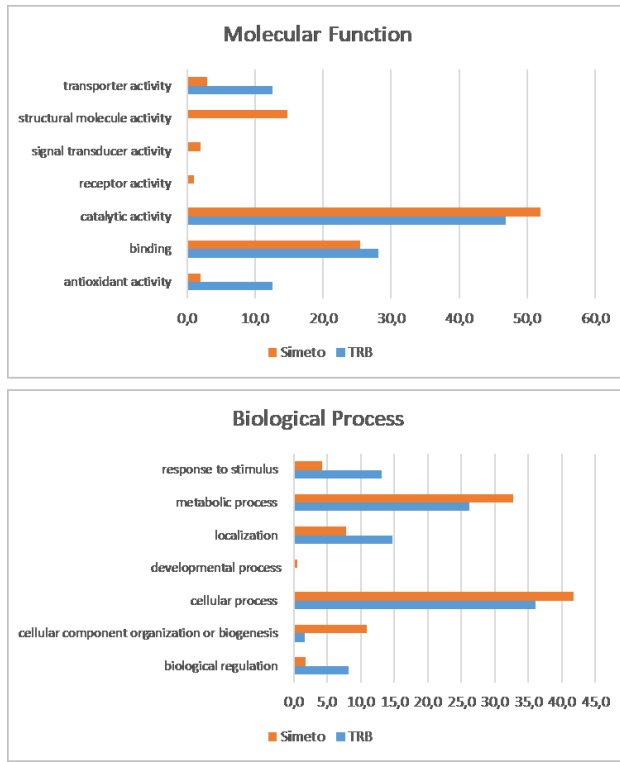


Figure 18. Gene Ontology of the unique proteins identified in the comparison *Timilia restre bianche* with *Simeto*

6.1.2 Qualitative comparison of the metabolic protein fractions of *Timilia restre bianche* and *Russello*

The list of proteins identified in the metabolic fraction of *Timilia restre bianche* and *Russello*, shared 349 proteins. On the contrary, 74 components were unique for the *cv Timilia* and 59 were exclusively found in the *cv Russello*.

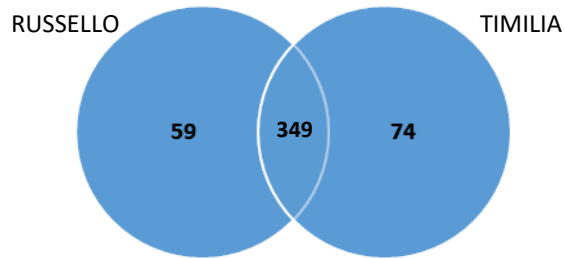


Figure 19. Venn diagram of the number of identified proteins in the metabolic fractions of cultivar *Timilia reste bianche* and *Russello*

The exclusive proteins of *Russello* and *Timilia* are mainly involved in the same Molecular Function: catalytic activity, binding and structural molecule activity. The signal trasducer (*receptor for activated C kinase 1A* and *Guanine nucleotide-binding protein subunit beta-like protein A*) and receptor activity (*V-type proton ATPase subunit B1*) are present only in the cultivar *Russello*. As above reported for the previous comparisons, the Biological Process distribution of unique proteins of *Russello* and *Timilia* does not show significant differences for almost all the biological processes reported (Fig. 20).

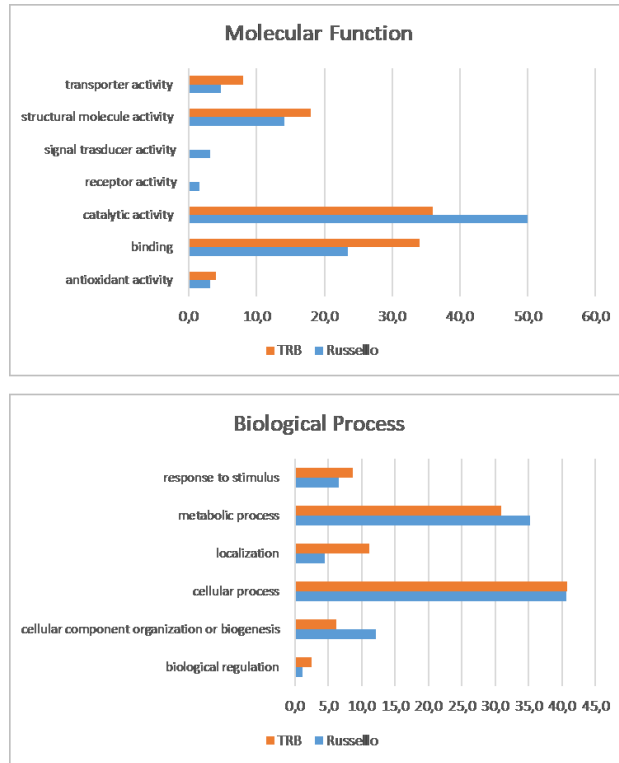


Figure 20. Gene Ontology of the unique proteins identified in the comparison *Russello* with *Timilia reste bianche*

6.1.3 Qualitative analysis of CM-like protein fractions of old and modern varieties

As expected, a selective separation of protein fractions was not achieved in the extraction method used. In fact, a cross contamination between the metabolic and CM-like fractions was observed. Therefore, a comparative analysis between the CM-like fraction of modern and ancient cultivars was conducted only considering the effective CM-like proteins. The CM-like enriched fractions contained mainly nine *alpha-amylase* and *alpha-amylase/trypsin inhibitors* (Tab. 13), together with minor amount of metabolic proteins. Comparison of these class of proteins revealed that all the investigated *cultivar* shows a very similar qualitative composition.

Table 13. CM-like proteins identified in the comparative analysis between old and modern *cultivar*

Acc. Number	Description
Q43723	Trypsin/alpha-amylase inhibitor CMX1/CMX3
Q43691	Trypsin/alpha-amylase inhibitor CMX2
P01885	Alpha-amylase inhibitor 0.19
P01083	Alpha-amylase inhibitor 0.28
P01084	Alpha-amylase inhibitor 0.53
P10846	Alpha-amylase inhibitor WDAI-3
P16159	Alpha-amylase/trypsin inhibitor CM16
P16851	Alpha-amylase/trypsin inhibitor CM2
P17314	Alpha-amylase/trypsin inhibitor CM3

6.2 Quantitative comparison

Three biological replicates of each *cultivar* were analyzed. The extraction procedure of the whole metabolic protein fraction was carried out one time for each biological replicate of each *cultivar*. Then, lysozyme was added as an internal standard in each extract and the solution was subjected to in-solution digestion followed by triplicate RP-nHPLC/nESI-MS/MS analysis and database search. Fig. 21 shows a scheme of the adopted procedure for each *cultivar*.

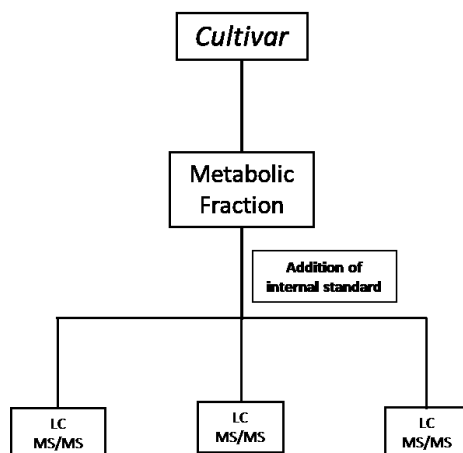


Figure 21. Scheme of the adopted procedure for each *cultivar*

The approach above described allowed the quantitative evaluation among the varieties investigated. Each old *cultivar* was compared with the modern *cultivar* *Simeto* and finally a comparison between the old genotypes was performed.

6.2.1 Quantitative comparison of the metabolic protein fractions between old *cultivar* and *Simeto*

In the comparison between *Russello vs Simeto* and *Timilia vs Simeto*, 34 and 46 proteins were found differentially expressed, respectively (Tab. 14-15). These proteins are displayed in the heat maps (Fig. 22-23). The relative protein abundance is represented in the map by a color and the map displays the expression trend of each protein in each sample. The protein abundance determined in the LC-MS replicate runs and in the biological replicates is reported in the heat map for each *cultivar*. *Simeto* was chosen as a reference. Over-expressed proteins in old *cultivar* are in the red zone of the map, instead the under-expressed proteins are in the green zone. On the left of the map there is a graph explaining the relationship between these proteins.

Table 14. List of proteins differentially expressed in the comparison between *Russello* and *Simeto*

Accession	Significance	Description	Fold change
P17314 IAAC3_WHEAT	200	Alpha-amylase/trypsin inhibitor CM3 OS=Triticum aestivum OX=4565 PE=1 SV=1	6.53
P82993 AMYB_HORVS	200	Beta-amylase OS=Hordeum vulgare subsp. spontaneum OX=77009 GN=BMY1 PE=1 SV=1	4.33
P16851 IAAC2_WHEAT	200	Alpha-amylase/trypsin inhibitor CM2 OS=Triticum aestivum OX=4565 PE=1 SV=2	11.51
P52589 PDI_WHEAT	200	Protein disulfide-isomerase OS=Triticum aestivum OX=4565 GN=PDI PE=2 SV=1	4.30
D2KFH1 AVLA4_WHEAT	200	Avenin-like a4 OS=Triticum aestivum OX=4565 PE=2 SV=1	25.92
P01084 IAA5_WHEAT	200	Alpha-amylase inhibitor 0.53 OS=Triticum aestivum OX=4565 PE=1 SV=1	3.37
P24296 NLTP1_WHEAT	200	Non-specific lipid-transfer protein (Fragment) OS=Triticum aestivum OX=4565 PE=1 SV=2	7.96
P0CZ08 AVLA3_WHEAT	200	Avenin-like a3 OS=Triticum aestivum OX=4565 PE=2 SV=1	27.19
P11143 HSP70_MAIZE	200	Heat shock 70 kDa protein OS=Zea mays OX=4577 GN=HSP70 PE=3 SV=2	4.58
P16347 IAAS_WHEAT	200	Endogenous alpha-amylase/subtilisin inhibitor OS=Triticum aestivum OX=4565 PE=1 SV=1	7.18
P37833 AATC_ORYSJ	200	Aspartate aminotransferase cytoplasmic OS=Oryza sativa subsp. japonica OX=39947 GN=Os01g0760600 PE=2 SV=1	3.63
P12810 HS16A_WHEAT	200	16.9 kDa class I heat shock protein 1 OS=Triticum aestivum OX=4565 GN=hsp16.9A PE=2 SV=1	6.42
Q9FRV0 CHIC_SECCE	200	Basic endochitinase C OS=Secale cereale OX=4550 GN=rscC PE=1 SV=1	5.30
Q10A30 ALFC2_ORYSJ	200	Fructose-bisphosphate aldolase 2 cytoplasmic OS=Oryza sativa subsp. japonica OX=39947 GN=FBA2 PE=2 SV=1	3.16
P82900 NLT2G_WHEAT	200	Non-specific lipid-transfer protein 2G OS=Triticum aestivum OX=4565 PE=1 SV=2	19.45
P30271 AMYB_SECCE	200	Beta-amylase (Fragment) OS=Secale cereale OX=4550 GN=BMY1 PE=2 SV=1	12.96
P62787 H4_MAIZE	200	Histone H4 OS=Zea mays OX=4577 GN=H4C7 PE=1 SV=2	8.45
P10846 IAA3_WHEAT	200	Alpha-amylase inhibitor WDAI-3 (Fragment) OS=Triticum aestivum OX=4565 GN=IHA-B1-2 PE=1 SV=1	4.70
P01543 THNB_WHEAT	200	Purothionin A-1 OS=Triticum aestivum OX=4565 GN=THI1.3 PE=1 SV=2	5.97
O64394 TRXH_WHEAT	200	Thioredoxin H-type OS=Triticum aestivum OX=4565 PE=2 SV=3	10.85
P10385 GLTA_WHEAT	200	Glutenin low molecular weight subunit OS=Triticum aestivum OX=4565 PE=3 SV=1	64.00
B8AL97 CUCIN_ORYSI	200	Cupincin OS=Oryza sativa subsp. indica OX=39946 GN=OsI_13867 PE=1 SV=1	6.04
Q43691 IACX2_WHEAT	200	Trypsin/alpha-amylase inhibitor CMX2 OS=Triticum aestivum OX=4565 PE=2 SV=1	4.09

Accession	Significance	Description	Fold change
P21569 CYPH_MAIZE	200	Peptidyl-prolyl cis-trans isomerase OS=Zea mays OX=4577 GN=CYP PE=2 SV=1	3.47
P23901 ALDR_HORVU	156,54	Aldose reductase OS=Hordeum vulgare OX=4513 PE=1 SV=1	3.13
P16159 IAC16_WHEAT	146,54	Alpha-amylase/trypsin inhibitor CM16 OS=Triticum aestivum OX=4565 PE=1 SV=1	3.00
Q75KH3 GRDH_ORYSJ	143,53	Glucose and ribitol dehydrogenase homolog OS=Oryza sativa subsp. japonica OX=39947 GN=Os05g0140800 PE=2 SV=2	2.96
P26517 G3PC1_HORVU	140,1	Glyceraldehyde-3-phosphate dehydrogenase 1 cytosolic OS=Hordeum vulgare OX=4513 GN=GAPC PE=2 SV=1	2.93
P08477 G3PC2_HORVU	135,82	Glyceraldehyde-3-phosphate dehydrogenase 2 cytosolic (Fragment) OS=Hordeum vulgare OX=4513 GN=GAPC PE=2 SV=1	3.33
Q948T6 LGUL_ORYSJ	82,89	Lactoylglutathione lyase OS=Oryza sativa subsp. japonica OX=39947 GN=GLYI-11 PE=1 SV=2	2.25
O64392 WHW1_WHEAT	82,48	Wheatwin-1 OS=Triticum aestivum OX=4565 GN=PR4A PE=1 SV=1	2.24
Q9ST57 SPZ2A_WHEAT	35,94	Serpin-Z2A OS=Triticum aestivum OX=4565 PE=1 SV=1	2.17
P82901 NLT2P_WHEAT	20,49	Non-specific lipid-transfer protein 2P OS=Triticum aestivum OX=4565 PE=1 SV=1	22.97

Table 15. List of proteins differentially expressed in the comparison between *Timilia reste bianche* and *Simeto*

Accession	Significance	Description	Fold change
P82993 AMYB_HORVS	200	Beta-amylase OS=Hordeum vulgare subsp. spontaneum OX=77009 GN=BMV1 PE=1 SV=1	5.94
P16159 IAC16_WHEAT	200	Alpha-amylase/trypsin inhibitor CM16 OS=Triticum aestivum OX=4565 PE=1 SV=1	3.28
P17314 IAAC3_WHEAT	200	Alpha-amylase/trypsin inhibitor CM3 OS=Triticum aestivum OX=4565 PE=1 SV=1	3.38
P16851 IAAC2_WHEAT	200	Alpha-amylase/trypsin inhibitor CM2 OS=Triticum aestivum OX=4565 PE=1 SV=2	10.41
P52589 PDI_WHEAT	200	Protein disulfide-isomerase OS=Triticum aestivum OX=4565 GN=PDI PE=2 SV=1	7.94
P01084 IAA5_WHEAT	200	Alpha-amylase inhibitor 0.53 OS=Triticum aestivum OX=4565 PE=1 SV=1	5.34
P26517 G3PC1_HORVU	200	Glyceraldehyde-3-phosphate dehydrogenase 1 cytosolic OS=Hordeum vulgare OX=4513 GN=GAPC PE=2 SV=1	3.48
D2KFH1 AVLA4_WHEAT	200	Avenin-like a4 OS=Triticum aestivum OX=4565 PE=2 SV=1	14.89
Q2A783 AVLB1_WHEAT	200	Avenin-like b1 OS=Triticum aestivum OX=4565 GN=AVNLB PE=1 SV=1	27.43

Accession	Significance	Description	Fold change
P42895 ENO2_MAIZE	200	Enolase 2 OS=Zea mays OX=4577 GN=ENO2 PE=2 SV=1	4.83
P08477 G3PC2_HORVU	200	Glyceraldehyde-3-phosphate dehydrogenase 2 cytosolic (Fragment) OS=Hordeum vulgare OX=4513 GN=GAPC PE=2 SV=1	6.04
P11143 HSP70_MAIZE	200	Heat shock 70 kDa protein OS=Zea mays OX=4577 GN=HSP70 PE=3 SV=2	4.53
Q42971 ENO_ORYSJ	200	Enolase OS=Oryza sativa subsp. japonica OX=39947 GN=ENO1 PE=1 SV=2	5.32
P24296 NLTP1_WHEAT	200	Non-specific lipid-transfer protein (Fragment) OS=Triticum aestivum OX=4565 PE=1 SV=2	8.00
P24067 BIP2_MAIZE	200	Luminal-binding protein 2 OS=Zea mays OX=4577 GN=BIPE2 PE=1 SV=3	3.71
P16347 IAAS_WHEAT	200	Endogenous alpha-amylase/subtilisin inhibitor OS=Triticum aestivum OX=4565 PE=1 SV=1	6.38
P12810 HS16A_WHEAT	200	16.9 kDa class I heat shock protein 1 OS=Triticum aestivum OX=4565 GN=hsp16.9A PE=2 SV=1	5.58
P37833 AATC_ORYSJ	200	Aspartate aminotransferase cytoplasmic OS=Oryza sativa subsp. japonica OX=39947 GN=Os01g0760600 PE=2 SV=1	4.64
P0CZ08 AVLA3_WHEAT	200	Avenin-like a3 OS=Triticum aestivum OX=4565 PE=2 SV=1	18.02
P23951 CHI2_HORVU	200	26 kDa endochitinase 2 OS=Hordeum vulgare OX=4513 PE=1 SV=1	3.38
P23901 ALDR_HORVU	200	Aldose reductase OS=Hordeum vulgare OX=4513 PE=1 SV=1	3.63
Q6F2Y7 CLPB1_ORYSJ	200	Chaperone protein ClpB1 OS=Oryza sativa subsp. japonica OX=39947 GN=CLPB1 PE=2 SV=1	3.86
Q43772 UGPA_HORVU	200	UTP--glucose-1-phosphate uridylyltransferase OS=Hordeum vulgare OX=4513 PE=2 SV=1	3.99
P12783 PGKY_WHEAT	200	Phosphoglycerate kinase cytosolic OS=Triticum aestivum OX=4565 PE=2 SV=1	10.10
P82901 NLT2P_WHEAT	200	Non-specific lipid-transfer protein 2P OS=Triticum aestivum OX=4565 PE=1 SV=1	22.89
Q10A30 ALFC2_ORYSJ	200	Fructose-bisphosphate aldolase 2 cytoplasmic OS=Oryza sativa subsp. japonica OX=39947 GN=FBA2 PE=2 SV=1	5.23
Q9FRV0 CHIC_SECCE	200	Basic endochitinase C OS=Secale cereale OX=4550 GN=rscC PE=1 SV=1	5.12
P82900 NLT2G_WHEAT	200	Non-specific lipid-transfer protein 2G OS=Triticum aestivum OX=4565 PE=1 SV=2	21.57
P30271 AMYB_SECCE	200	Beta-amylase (Fragment) OS=Secale cereale OX=4550 GN=BMY1 PE=2 SV=1	12.95
P62787 H4_MAIZE	200	Histone H4 OS=Zea mays OX=4577 GN=H4C7 PE=1 SV=2	9.27
P01083 IAA2_WHEAT	200	Alpha-amylase inhibitor 0.28 OS=Triticum aestivum OX=4565 GN=IMA1 PE=1 SV=3	64.00
P10846 IAA3_WHEAT	200	Alpha-amylase inhibitor WDAI-3 (Fragment) OS=Triticum aestivum OX=4565 GN=IHA-B1-2 PE=1 SV=1	3.36

Accession	Significance	Description	Fold change
P02276 H2A2_WHEAT	200	Histone H2A.2.1 OS=Triticum aestivum OX=4565 PE=1 SV=1	5.04
P01543 THNB_WHEAT	200	Purothionin A-1 OS=Triticum aestivum OX=4565 GN=THI1.3 PE=1 SV=2	3.49
O64394 TRXH_WHEAT	200	Thioredoxin H-type OS=Triticum aestivum OX=4565 PE=2 SV=3	7.89
B8AL97 CUCIN_ORYSI	200	Cupincin OS=Oryza sativa subsp. indica OX=39946 GN=OsI_13867 PE=1 SV=1	3.98
Q65XA0 DHAR1_ORYSJ	200	Probable glutathione S-transferase DHAR1 cytosolic OS=Oryza sativa subsp. japonica OX=39947 GN=DHAR1 PE=1 SV=1	3.90
Q43691 IACX2_WHEAT	200	Trypsin/alpha-amylase inhibitor CMX2 OS=Triticum aestivum OX=4565 PE=2 SV=1	4.93
Q07661 NDK1_ORYSJ	200	Nucleoside diphosphate kinase 1 OS=Oryza sativa subsp. japonica OX=39947 GN=NDKR PE=1 SV=1	4.07
P21569 CYPH_MAIZE	200	Peptidyl-prolyl cis-trans isomerase OS=Zea mays OX=4577 GN=CYP PE=2 SV=1	3.89
P27806 H1_WHEAT	200	Histone H1 OS=Triticum aestivum OX=4565 PE=2 SV=2	6.25
Q75KH3 GRDH_ORYSJ	156,54	Glucose and ribitol dehydrogenase homolog OS=Oryza sativa subsp. japonica OX=39947 GN=Os05g0140800 PE=2 SV=2	3.11
P04568 EM1_WHEAT	147,5	Em protein OS=Triticum aestivum OX=4565 GN=EM PE=2 SV=1	2.01
Q948T6 LGUL_ORYSJ	105,18	Lactoylglutathione lyase OS=Oryza sativa subsp. japonica OX=39947 GN=GLYI-11 PE=1 SV=2	2.69
Q9XHS0 RS12_HORVU	98,81	40S ribosomal protein S12 OS=Hordeum vulgare OX=4513 GN=RPS12 PE=2 SV=1	9.90
P21641 OLEO3_MAIZE	20,45	Oleosin Zm-II OS=Zea mays OX=4577 GN=OLE18 PE=1 SV=1	3.55

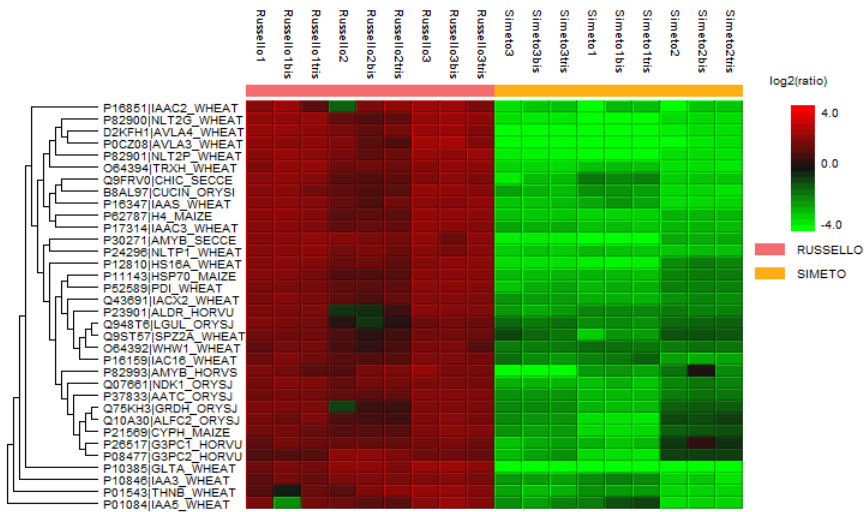


Figure 22. Heat map of the quantitative comparison between *Russello* and *Simeto*

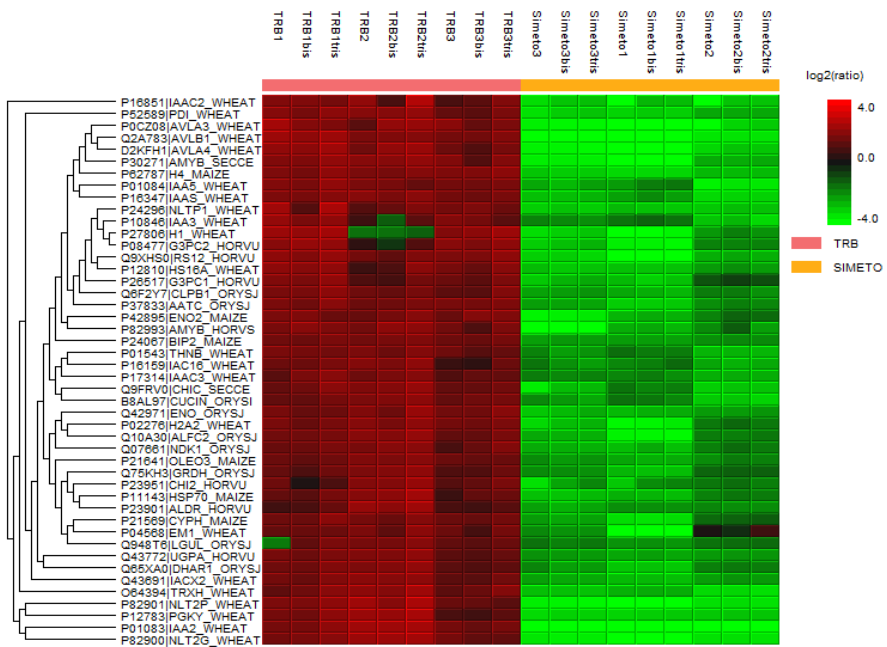


Figure 23. Heat map of the quantitative comparison between *Timilia reste bianche* and *Simeto*

The quantitative evaluation of the identified proteins detected that about 70% of the differentially expressed proteins are common to both old *cultivar*. As described before, proteins contained in the albumin and globulin fractions (metabolic proteins) are responsible for wheat allergies, and some of the main allergenic proteins of these fractions were found over expressed in the old genotypes, such as: *alpha-amylase/trypsin inhibitor CM3*,^{9,36} *CM2*^{9,37} and *CM16*,^{9,36} *alpha-amylase inhibitor 0.53*^{9,37} and *WDAI-3*,⁹ *trypsin/alpha-amylase inhibitor CMX2*,⁹ *non-specific lipid-transfer protein 2G* and *2P*,^{9,38,39} *thioredoxin H-type*,⁹ *beta-amylase*,^{9,37} *endogenous alpha-amylase/subtilisin inhibitor*³⁷ and *fructose-bisphosphate aldolase 2 cytoplasmic*.⁹ Moreover, in the comparison of *Russello* vs *Simeto* another allergenic protein was found: *serpin-Z2A*.^{9,40}

The others not allergenic proteins common to the two old cultivars are mainly involved in Molecular Function (binding and catalytic activity) and in Biological Process (biological regulation, cellular process, developmental process, metabolic process and response to stimulus). In details, *Avenin-like a4* is involved in a specific Molecular Function, nutrient reservoir activity and *Protein disulphide-isomerase* is involved in “cell redox homeostasis” (biological process), that is any process that maintains the redox environment of a cell or compartment within a cell.

Wheatwin-1, a unique protein identified in *Russello*, shows the same activity above described.

Avenin-like b1, *Enolase 2*, *Enolase*, *Luminal-binding protein 2*, *Avenin-like a3*, *26KDa endochitinase 2*, *Histone H2A.2.1*, *Probable glutathione S-transferase DHAR1*, *Histone H1*, *Em protein*, *40S ribosomal protein S12* and *Oleosin Zm-II*, are unique proteins of *Timilia*. These proteins are involved in the same activity listed above, with only one exception for *Probable glutathione S-transferase DHAR1* that is involved in “cellular oxidant detoxification”. This Cellular Process is part of the defence response of the plant and it participates in the elimination of the toxic superoxide radicals or hydrogen peroxide.

6.2.2 Quantitative comparison of the metabolic protein fractions between old cultivar

In the comparison between *Timilia* vs *Russello* the second one was chosen like a reference. 8 and 7 proteins were found over-expressed and under expressed, respectively (Tab. 16). These proteins are displayed in the heat map in Fig. 24.

Only two allergenic proteins were found in the group of differentially expressed proteins. In particular, *serpin-Z2A* and *alpha-amylase inhibitor 0.28* are particularly over-expressed in the cultivar *Russello* and *Timilia*, respectively.

The under expressed proteins in *Timilia*: *26 kDa endochitinase 2*, *Chaperone protein ClpB1*, *Coleoptile phototropism protein 1*, *Wheatwin-1* and *Succinate--CoA ligase [ADP-forming] subunit beta mitochondrial*, are mainly involved in catalytic activity and metabolic processes of the plant. In particular, *26 kDa endochitinase 2* and *Wheatwin-1* show, also, an activity in the defence response of this variety. They are involved in the restriction of damage to the organism attacked or prevention/recovery from the infection caused by the attack. At the end, two *Glutenin low molecular weight subunit* are found in this group.

Avenin-like b1, *Subtilisin-chymotrypsin inhibitor WSCI*, *Defensin Tk-AMP-D4*, *Sucrose synthase 2*, *Sucrose synthase 1* and *60S ribosomal protein L7a-1*, are found over-expressed in *Timilia*. These proteins are involved in Molecular Function (binding and catalytic activity) and in Biological Process (cellular process, metabolic process and cellular component organization or biogenesis). In particular: *Avenin-like b1* is a seed storage protein involved in the storage of nutritious substrates, *Subtilisin-chymotrypsin inhibitor WSCI* participates in any process that results in a change in state or activity of a cell or an organism (in terms of movement, secretion, enzyme production, gene expression, etc.) as a result of a stimulus indicating damage to the organism and *Defensin Tk-AMP-D4* shows a reaction against the presence of a fungus in order to protect the cell or organism.

Table 16. List of proteins differentially expressed in the comparison between *Timilia reste bianche* and *Russello*

Accession	Significance	Description	Fold change
Q2A783 AVLB1_WHEAT	200	Avenin-like b1 OS=Triticum aestivum OX=4565 GN=AVNLB PE=1 SV=1	56.42
Q9ST57 SPZ2A_WHEAT	200	Serpin-Z2A OS=Triticum aestivum OX=4565 PE=1 SV=1	0.04
P01083 IAA2_WHEAT	200	Alpha-amylase inhibitor 0.28 OS=Triticum aestivum OX=4565 GN=IMA1 PE=1 SV=3	64.00
P10386 GLTB_WHEAT	200	Glutenin low molecular weight subunit 1D1 OS=Triticum aestivum OX=4565 PE=2 SV=1	0.04
P10385 GLTA_WHEAT	200	Glutenin low molecular weight subunit OS=Triticum aestivum OX=4565 PE=3 SV=1	0.18
P82977 ICIW_WHEAT	200	Subtilisin-chymotrypsin inhibitor WSCI OS=Triticum aestivum OX=4565 PE=1 SV=2	4.95
P84971 DEF4_TRIKH	109,32	Defensin Tk-AMP-D4 OS=Triticum kiharae OX=376535 PE=1 SV=1	3.01
P23951 CHI2_HORVU	105,46	26 kDa endochitinase 2 OS=Hordeum vulgare OX=4513 PE=1 SV=1	0.40
P31923 SUS2_HORVU	86,96	Sucrose synthase 2 OS=Hordeum vulgare OX=4513 GN=SS2 PE=1 SV=1	2.30
P35685 RL7A1_ORYSJ	79,85	60S ribosomal protein L7a-1 OS=Oryza sativa subsp. japonica OX=39947 GN=RPL7A-1 PE=2 SV=1	2.21
Q6F2Y7 CLPB1_ORYSJ	77,33	Chaperone protein ClpB1 OS=Oryza sativa subsp. japonica OX=39947 GN=CLPB1 PE=2 SV=1	0.46
P31922 SUS1_HORVU	64,8	Sucrose synthase 1 OS=Hordeum vulgare OX=4513 GN=SS1 PE=1 SV=1	2.10
Q5KS50 NPH3_ORYSJ	41,33	Coleoptile phototropism protein 1 OS=Oryza sativa subsp. japonica OX=39947 GN=CPT1 PE=2 SV=1	0.05
O64392 WHW1_WHEAT	39,58	Wheatwin-1 OS=Triticum aestivum OX=4565 GN=PR4A PE=1 SV=1	0.28
Q6K9N6 SUCB_ORYSJ	39,06	Succinate--CoA ligase [ADP-forming] subunit beta mitochondrial OS=Oryza sativa subsp. japonica OX=39947 GN=Os02g0621700 PE=1 SV=1	0.47



Figure 24. Heat map of the quantitative comparison between *Timilia reste bianche* and *Russello*

7. A manually curated database of metabolic proteins from *Triticum Aestivum*

7.1 Introduction

As describe above, wheat plays an important role in many adverse reactions. The albumin/globulin fraction includes the main important allergenic proteins correlated with wheat allergy.

Mass spectrometry is a technique alternative to immune based methods for the identification of allergenic determinant. The immunoreactive protein is digested using trypsin and analysed by mass spectrometry. The identification is performed by bioinformatics search on a repository of protein sequences, such as Uniprot. Uniprot is constituted of two main sections: swissprot, including reviewed sequences and tremble, including unreview sequences. An unreview sequence may be a fragment or a repetition of a reviewed sequence.

Central to the mass spectrometry based approach is the requirement for a well curated set of sequence from other sources such as UniProt. In general, the area of plant proteomics suffers for the lack of curated sequence database. Out of ~ 35 000 cultivated plant species, only 37 have sequenced and functionally annotated genomes.⁴¹

In UniProt (release May 2018), the reviewed *Viridiplantae* database contains 39.456 sequences, but UniProt has focused only on the curation of plant protein sequences from *Arabidopsis thaliana* and *Oryza sativa*. Furthermore, in the plant area there is an additional problem: genomic sequencing is very difficult due to the large size and complexity of the plant genomes, because several species, including major crops such as wheat (*triticum*) are polyploid. This problem has motivated many researchers to develop manually curated database to support protein identification.⁴¹

The aim of this part of the work was to develop a manually curated database, in FASTA format, of metabolic proteins, focused exclusively on hexaploid bread wheat (*Triticum aestivum*).

7.2 Materials and method

7.2.1 Identification of wheat seed protein sequences accession in UniProt

Accession numbers were obtained from Louise J. Salt et al., 2004. This paper contains a list of wheat seed metabolic proteins. Some of these proteins were identified by homology with other species, such as *Oryza sativa*, *Hordeum vulgare*, *Secale cereale*, *Zea mays*, *H. spontaneum* and *Lolium perenne*. Today, these sequences of *Triticum aestivum* are included in UniProt. This list was used to create a verified set of seed proteins to start the creation of the database (Tab. 17).

7.2.2 Database construction and curation

In Table 17, are reported the description, accession number, protein family classification (Pfam name), gene location, allergen name and reference for each protein. Also, the description contains information about the characterization and the status in UniProt of the protein. The status of UniProt contains information about the evidence of the proteins (experimental evidence at protein level, protein inferred from homology, experimental evidence at transcript level, etc.) and the annotation score. The gene location and the allergen name were found using <https://plants.ensembl.org> and UniProt, respectively. An independent database set was developed for each family protein of this and subsequently the data bases were merged into a single database (Fig. 25).

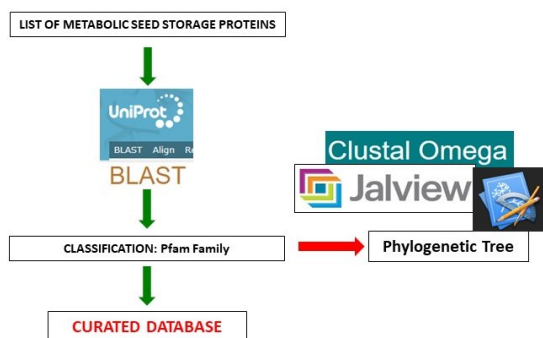


Fig 25. Scheme of the adopted procedure

The UniProt accession for each verified sequence was BLAST (basic logic algorithm search tool) searched against the entire UniProt database, and the search parameters set to show 1000 results. For each BLAST search a single file per family was saved. From the extended list, we only took into consideration protein sequences of *T. aestivum* showing an identity greater than 70%. Redundant sequences were manually removed. Each sequence shortlisted was further BLAST searched against the entire UniProt, as described above, to verify the presence of redundancies. Selected sequences were aligned using the Clustal Omega software and saved as a MSF file. These files were viewed in Jalview to observe multiple sequence alignment to show regions of high homology. At this point, the sequences were manually interrogated, to remove incomplete and incorrectly annotated sequences. For each subgroup a set of unique and full length sequences remained. The resulting curated sequence sets were combined to develop the complete database. The final list of the verified UniProt sequence accessions can be found in the Table 18.

7.2.3 Phylogenetic analysis

Phylogenetic analysis was carried out using the BLOSUM62 algorithm in JalView for each Pfam and the resulting tree viewed and manipulated in FigTree. These were used to illustrate the relationships within the subgroup database.

7.3 Result and discussion

7.3.1 Database development and characterization

Verified sequence representing the proteins in Table 17 were used to control the unreviewed and reviewed UniProt database. 579 sequences of *T. aestivum* were downloaded and subsequently 31% of redundant sequences were removed in the initial step (Fig. 26).

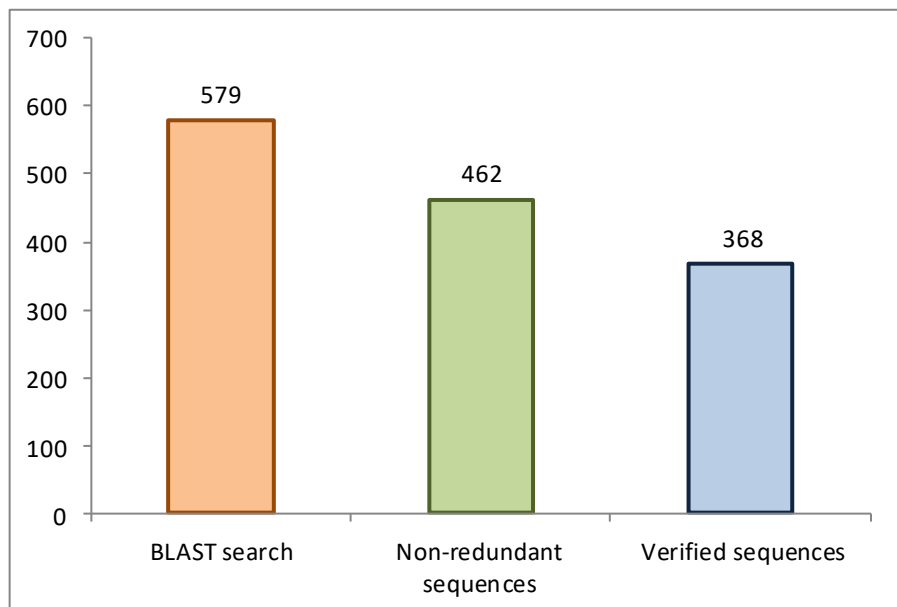


Fig 26. Database composition

For example, UniProt sequence #Q53YX8 was removed from the α -*amylase/trypsin inhibitor* subunit sequence set because its sequence is the same of the UniProt sequence #P17314. The UniProt sequence #B5B0D5 was removed because the difference with the sequence #P16159 is a single aminoacidic substitution occurring in the signal peptide. The final database was composed of 460 sequences (Tab. 19), in which only 368 sequences contain a reference in UniProt. These sequences were classified by using the

Pfam classification in UniProt. The database composition is reported in Table 18. Phylogenetic analysis of the aligned sequences from database revealed a clustering based on the Pfam classification (Fig. 27). The phylogenetic tree of the complete curated database was created by using the algorithm in Jalview and then manipulated in FigTree. For example, the Pfam “tryo-alpha-amyl” contains all *alpha amylase/trypsin inhibitors*. A detailed analysis shows clustering of these proteins into four groups corresponding to the *alpha-amylase inhibitor 0.28*, *alpha-amylase inhibitor 0.19-0.53*, *alpha-amylase/trypsin inhibitors CM1-CM2-CM3-CM16* and *trypsin/alpha-amylase inhibitors CMX1/CMX3-CMX2*. Some of these groups included more inhibitors, because they shared the same set of proteins (Fig. 28). The phylogenetic tree displays the distribution of these sequences on different nodes, but the average sequence identity is very high (90.5%) and they belong to the same cluster. When different set of proteins are classified in UniProt by using the same Pfam name, these proteins are grouped in the same cluster. One example is the proteins set of *embryo specific protein* and *lipoprotein-like*. There is only one exception, the set proteins of *dehydroascorbate reductase* and *glutathione transferase* which in UniProt have the same Pfam name, whereas in the phylogenetic tree these sequences are not grouped.

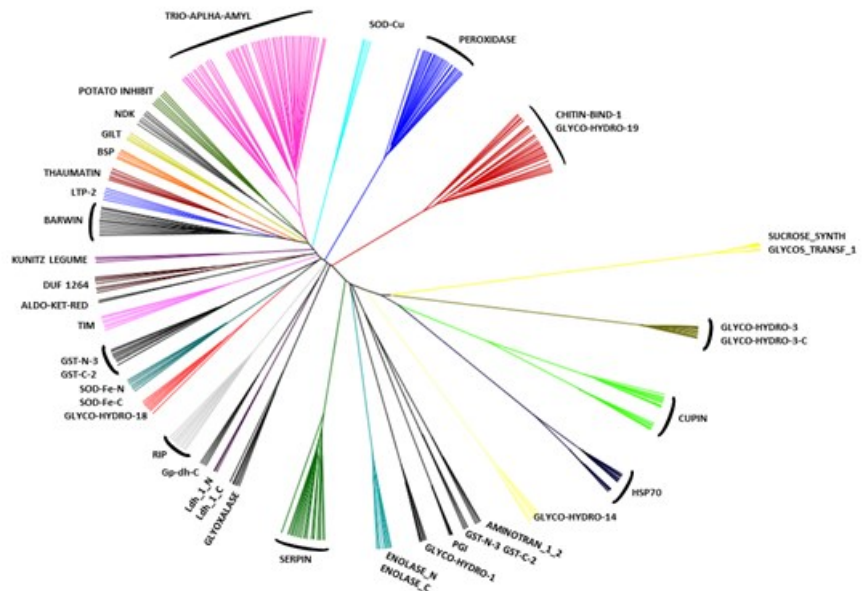


Fig 27. Phylogenetic tree for the complete curated database

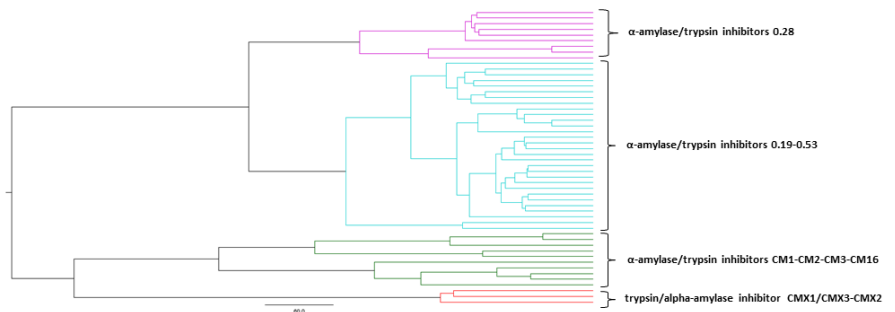


Fig. 28. A phylogenetic analysis of the “tryo-alpha-amyl” sequences in the database

8. Conclusion

In this work, a comparative proteomic analysis of the metabolic fraction, of three different durum wheat varieties (*Simeto*, *Russello* and *Timilia reste bianche*) was performed at qualitative and quantitative levels. *Simeto* is representative of the commercial *cultivar* used in the current commercial practice. Instead, *Russello* and *Timilia* are old Sicilian durum wheat landraces and they are traditionally cultivated in some areas of Sicily as Agrigento, Caltanissetta, Ragusa and Trapani. They are used in the production of bread and pasta with high digestibility.

The qualitative proteomic analysis of these durum wheat varieties revealed a remarkable similarity in the protein composition between old and modern *cultivar*. In particular, comparison of the proteins identified in the metabolic fraction of the *cultivar* investigated revealed that most of them are involved in binding and catalytic activity and play a role in the same biological processes. The qualitative analysis of the CM-like proteins revealed that the same proteins were identified in the varieties investigated. In conclusion, relevant differences were not found at qualitative level.

Moreover, also a quantitative evaluation was performed and this analysis allowed to ascertain that some components are differentially expressed in these genotypes. In the comparison between *Russello* and *Timilia vs Simeto* some of the main allergenic metabolic proteins were found over expressed in the old genotypes (*alpha-amylase/trypsin inhibitor CM3-CM2-CM16*, *alpha-amylase inhibitor 0.53- WDAI-3*, *non-specific lipid-transfer protein 2G* and *2P*, *thioredoxin H-type*, *beta-amylase*, *endogenous alpha-amylase/subtilisin inhibitor* and *fructose-bisphosphate aldolase 2 cytoplasmic*). Moreover, in the comparison of *Russello vs Simeto* another allergenic protein, *serpin-Z2A*, was found exclusively in *Russello*. Only two allergenic proteins were found differentially expressed in the comparison of *Timilia vs Russello*. In particular, *serpin-Z2A* and *alpha-amylase inhibitor 0.28* are particularly over-expressed in the cultivar *Russello* and *Timilia*, respectively.

One of the main aim of many proteomics analysis is to identify the largest number of proteins with the highest quality. In wheat, the possibility of complete protein identification is hampered by the lack of information about the full genome sequence. UniProt has focused only on the curation of plant protein sequences from *Arabidopsis thaliana* and *Oryza sativa*. This problem has motivated many researchers to develop manually curated databases to support protein identification. These curated databases can be used in the proteomic analysis for identification purposes. Future work will focus on the extension this database to include other wheat species such as *Triticum durum*.

In conclusion, this study represents the first molecular characterization of the metabolic and CM-like protein fractions at qualitative and quantitative levels of the old Sicilian landraces *Russello* and *Timilia reste bianche*. This comparison would help to understand the relationship occurring between protein profile of old wheat varieties and potential benefits for human consumption.

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Figure 18. Gene Ontology of the unique proteins identified in the comparison *Timilia reste bianche* with *Simeto*

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11. List of Publications

1. Gallina S., V. Cunsolo, R. Saletti, V. Muccilli, A. Di Francesco, S. Foti, A. M. Lorenzten, P. Roepstorff. (2016). "Sequence characterization and glycosylation sites identification of donkey milk lactoferrin by multiple enzyme digestions and mass spectrometry". *Amino Acids* 48 (7): 1569-1580.
2. Cunsolo V., E. Fasoli, A. Di Francesco, R. Saletti, V. Muccilli, S. Gallina, P. G. Righetti, S. Foti. (2016). "Polyphemus, Odysseus and the Ovine milk proteome". *Journal of Proteomics*: 5874.
3. Cunsolo V., R. Saletti, V. Muccilli, S. Gallina, A. Di Francesco, S. Foti. (2017). "Proteins and bioactive peptides from donkey milk: The molecular basis for its reduced allergenic properties". *Food Research International* 99: 41-57.
4. Criscione A., V. Cunsolo, S. Tumino, A. Di Francesco, S. Bordonaro, V. Muccilli, R. Saletti, D. Marletta. (2018). "Polymorphism at donkey β -lactoglobulin II locus: identification and characterization of a new genetic variant with a very low expression". *Amino Acids* 50 (6): 735-746.

The results of my PhD activity will be published in three papers in preparation, dealing with:

- Proteomic analysis of the metabolic fractions in modern and old wheat genotypes: a qualitative comparison;
- Proteomic analysis of the metabolic fractions in modern and old wheat genotypes: a quantitative comparison;
- A manually curated database of metabolic proteins from *Triticum Aestivum*.

12. Conference Contributions

1. S. Gallina, R. Saletti, V. Cunsolo, V. Muccilli, A. Di Francesco, S. Foti, P. Roepstorff. "High resolution mass spectrometry in the analysis of site-specific donkey milk lactoferrin glycosylation" - XXXVI Organic Division 13-17 September 2015, Bologna. Abstract book pag. 207.
2. V. Cunsolo, V. Muccilli, R. Saletti, S. Gallina, A. Di Francesco, S. Foti. "Integrated bottomup/top-down ms-based approach for sequence determination of a donkey' CSN1S2 duplicate gene products" - XXXVI Organic Division, 13-17 September 2015, Bologna. Abstract book pag. 208.
3. A. Di Francesco, V. Cunsolo, E. Fasoli, V. Muccilli, M. Nicoletti, S. Gallina, R. Saletti, P. G. Righetti, S. Foti. "Sheep's Milk: a deep exploration of the low-abundance proteome" - XI Annual ItPA Conference, 16-19 May 2016, Perugia. Abstract book pag. 53.
4. A. Di Francesco, R. Saletti, V. Cunsolo, V. Muccilli, P. De Vita, S. Gallina, S. Foti. "Comparative proteomic analysis of the metabolic fractions of Russello and Simeto durum wheat varieties" – Workshop SCI 2016-17, 9-10 February 2017, Messina. Abstract book P-8.
5. A. Di Francesco, R. Saletti, V. Cunsolo, V. Muccilli, P. De Vita, S. Gallina, S. Foti. "Ancient and modern durum wheat varieties: a comparative proteomic analysis of the metabolic fractions" - XII Annual ItPA Conference, 12-15 June, 2017, Lecce. Abstract book pag. 59. (Poster Prize)
6. Antonella Di Francesco, Rosaria Saletti, Vincenzo Cunsolo, Vera Muccilli, Pasquale De Vita and Salvatore Foti. "Analisi proteomica comparativa della frazione metabolica di varietà antiche e moderne di grano duro". 11° AISTEC Congress, 22-24 November 2017, Roma. Abstract book pag. 193.
7. Vincenzo Cunsolo, Antonella Di Francesco, Rosaria Saletti, Vera Muccilli, Salvatore Foti. "Grani antichi e grani moderni: caratterizzazione della frazione proteica mediante gli approcci proteomici". 11° AISTEC Congress, 22-24 November 2017, Roma. Abstract book pag. 41.

8. A. Di Francesco, V. Cunsolo, R. Saletti, V. Muccilli, P. De Vita, S. Foti.
"Grani antichi e grani moderni: analisi proteomica comparative della frazione metabolica". Congresso congiunto delle sezioni Sicilia e Calabria SCI 2018, 9-10 Febbraio 2018, Catania. Abstract book pag. 58.
9. A. Di Francesco, V. Cunsolo, R. Saletti, V. Muccilli, P. De Vita, S. Foti.
"Metabolic fractions of ancient and modern durum wheat varieties: a comparative proteomic analysis" - 36th INFORMAL MEETING ON MASS SPECTROMETRY, 6-9 May, Koszeg. Abstract book pag. 101.
10. A. Di Francesco, R. Saletti, V. Cunsolo, V. Muccilli, P. De Vita, S. Foti.
"Comparative proteomic analysis of the metabolic fractions composition in modern and old wheat genotypes". XXII International Mass Spectrometry Conference, 26-31 August 2018, Firenze. Abstract book pag. 873.
11. Vincenzo Cunsolo, Michele A. De Santis, Antonella Di Francesco, Rosaria Saletti, Marcella M. Giuliani, Zina Flagella, Salvatore Foti. "Investigation of the gliadin fraction composition in old and modern wheat genotypes". XXII International Mass Spectrometry Conference, 26-31 August 2018, Firenze. Abstract book pag. 1119.
12. A. Di Francesco, R. Saletti, V. Cunsolo, V. Muccilli, P. De Vita, S. Foti.
"Metabolic fractions of old and modern durum wheat varieties: a comparative proteomic analysis". XXIII Annual ItPA Conference, 5-7 September, Como. Abstract book pag. 35.
13. Vincenzo Cunsolo, Michele A. De Santis, Antonella Di Francesco, Rosaria Saletti, Marcella M. Giuliani, Zina Flagella, Salvatore Foti. "Comparison of the gliadin fraction between old and modern durum wheat genotypes: a proteomic approach". XXIII Annual ItPA Conference, 5-7 September, Como. Abstract book pag. 66.

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