GENETIC AND GENOMIC APPROACHES TO INVESTIGATE ZYMV RESISTANCE IN *Cucurbita pepo*
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SOMMARIO


È importante individuare la corretta locazione sul genoma dei geni di resistenza caratterizzati, affinché si possano individuare marcatori molecolari ad essi associati da impiegare nelle successive fasi di selezione genetica.

Il principale obiettivo di questo lavoro è l’identificazione di marcatori molecolari associati ai geni noti per conferire resistenza a ZYMV, che potrebbero risultare utili per il miglioramento genetico di varietà di zucchino tra cui la varietà locale ‘San Pasquale’ tipica della regione Campania. Il secondo obiettivo di questo lavoro è rappresentato dall’individuazione di nuovi geni candidati, e relativi marcatori SNP ad essi associati, coinvolti nel meccanismo di resistenza a ZYMV in _Cucurbita pepo_ mediante un approccio trascrittomico. L’ultimo obiettivo è quello di effettuare una predizione genica dei geni di resistenza R nel genoma di zucchino.

In primo luogo, sono state ottenute due popolazioni segreganti F₂ dall’incrocio di varietà di zucchino suscettibili (‘True French’ e ‘San Pasquale’) con una varietà resistente al virus (‘381e’) e tre BC₁ dal successivo reincrocio con la ‘San Pasquale’. Le valutazioni fenotipiche dei sintomi effettuate sulle 5 popolazioni sono in accordo con i rapporti di segregazione proposti da Paris e Cohen (2000) per 3 geni complementari dominanti sono candidati per la resistenza a ZYMV.

I dati provenienti dall’uso di una piattaforma SNP (Illumina GoldenGate; Esteras _et al._, 2012) hanno consentito di valutare la distanza genetica delle linee ‘True French’ (TFR) ‘381e’ e ‘San Pasquale’ (SPQ) e di individuare 6 polimorfismi SNP tra le accessioni suscettibili e quelle resistenti al virus. Questi SNP sono stati valutati sui parentali e le relative popolazioni segreganti, già caratterizzate fenotipicamente per la resistenza al virus. Le analisi molecolari HRM, hanno evidenziato tre marcatori SNP putativamente associati ai geni di resistenza (Zym-1, Zym-2 e Zym-3) proposti da Paris e Cohen (2000). I risultati dell’analisi di segregazione sono in accordo con il rapporto atteso di 3 resistenti su 1 suscettibile ($\chi^2 P=0,8$). Tale rapporto è stato dedotto ipotizzando che il gene Zym-1 sia il gene maggiore rispetto ai geni minori Zym-2 e Zym-3, ipotesi confermata.
anche dalla percentuale di co-segregazione calcolata tra il genotipo e il fenotipo di ogni singola pianta e dalla percentuale di co-segregazione tra i singoli marcatori testati ed il fenotipo. Il test del chi-quadrato suggerisce che esiste una forte relazione tra lo SNP1 ed il gene maggiore Zym-1. Anche la valutazione molecolare effettuata sulla popolazione F2 ‘TFRx381e’, attraverso un’analisi CAPS, ha confermato l’associazione tra lo SNP1 ed il gene maggiore Zym-1 (χ2 P=0,7) Inoltre, il test di correlazione Mantel condotto sui dati genotipici e fenotipici delle due popolazioni segreganti F2 ‘TFRx381e’ e ‘SPQx381e’ ha mostrato che esiste una forte correlazione tra le due matrici in esame (r(AB) = 0,112).

Al fine di identificare nuovi geni candidati e nuovi marcatori SNP associati alla resistenza a ZYMV è stato effettuato un esperimento di sequenziamento del trascrittoma (RNA-seq) di piante inoculate del genotipo suscettibile ‘TFR’ e del genotipo resistente ‘381e’. L’analisi dei geni differenzialmente espressi (DEG) ha evidenziato 206 geni comuni nelle due linee isogeniche analizzate, 1956 espressi esclusivamente nel genotipo suscettibile e 1006 trascritti nel genotipo resistente a ZYMV.

Al fine di ottenere una classificazione in categorie funzionali ‘Gene Onthology’ (GO) è stata effettuata l’annotazione funzionale tramite il software Blast2GO (B2GO; Conesa et al., 2005) del trascrittoma di riferimento. L’analisi ha rivelato che il 36,54 % dei trascritti è ascritto in almeno una delle tre categorie funzionali GO (molecular function, biological process e cellular component). Sulla base di questa analisi è stato possibile annotare i trascritti differenzialmente espressi nei due genotipi. L’annotazione funzionale dei DEG nei genotipi analizzati ha permesso di identificare, una relativa abbondanza di trascritti che codificano per domini LRR e Serina/Treonina kinasi in entrambi i genotipi, mentre una più alta concentrazione di trascritti codificanti Elicasi ATP-dipendenti è stata riscontrata nel genotipo resistente.

I dati provenienti dal sequenziamento del trascrittoma di ‘TFR’ e ‘381e’ sono stati impiegati anche per effettuare un’analisi di SNP calling nei confronti del genoma di riferimento. Tale analisi ha permesso di evidenziare 3357 e 14327 SNP unici caratterizzanti rispettivamente ‘TFR’ e ‘381e’. Gli SNP1 e SNP2, putativamente correlati al gene Zym-1 e Zym-2, sono stati identificati tra i 14327 SNPs del genotipo ‘381e’, mentre lo SNP3 nei 3357 SNP unici del genotipo ‘TFR’.

Dal confronto delle liste dei geni contenenti gli SNP e le liste dei DEG è stato possibile identificare i geni regolati in modo positivo o negativo, espressi nei due genotipi in relazione all’attacco del virus e polimorfici rispetto al genoma di riferimento. Tra i geni differenzialmente espressi e polimorfici è stata evidenziata un’alta percentuale di geni codificanti per ‘Protein Kinase’, ‘Chloroplast Protein Kinase’ e ‘Leucine Reach Repeats’ in entrambe i genotipi oltre ad una rilevante quantità di altre proteine come Taumatina, Ubiquitina, Fosfatasi, ed ormoni quali Auxine ed Etilene, tipicamente coinvolti nelle reazioni di risposta della pianta a stress biotici.
Al fine di identificare gli SNP unici dei due genotipi correlati a geni di resistenza, è stato interrogato il proteoma di *Cucurbita pepo* con un predittore specifico per geni di resistenza chiamato MATRIX R (Hermoso Pulido *et al.*, 2014). I risultati hanno evidenziato che dei 3357 e 14327 SNP unici dei genotipi ‘TFR’ and ‘381e’, 251 SNP e 1895 SNP, rispettivamente, ricadono in geni o domini tipici coinvolti in meccanismi di difesa della pianta. In particolare, 100 SNP del genotipo ‘381e’ ricadono in soli 3 geni R localizzati sullo scaffold2494 in cui è presente anche lo SNP1 associato al gene maggiore *Zym-1*, così come 19 dei 251 SNP identificati nel genotipo ‘TFR’ ricadono su in 3 geni R dello scaffold1174 in cui risiede lo SNP3 associato al gene minore *Zym-3*.

Concludendo, lo studio dell’ereditarietà della resistenza a ZYMV attraverso le analisi fenotipiche e genotipiche, condotte su cinque popolazioni segreganti, hanno messo in evidenza il coinvolgimento di un gene principale e di due geni minori nel meccanismi di resistenza a tale virus. I saggi *in vivo* e le analisi molecolari hanno permesso di mostrare una forte relazione tra i tre marcatori SNP identificati e questi geni di resistenza, consentendo di avere uno strumento molecolare utile per una selezione più mirata. Le analisi trascrittomiche condotte su due linee isogeniche ‘True French’ e ‘381e’, hanno permesso di identificare geni R polimorfi tra il genotipo resistente e quello suscettibile espressi in modo diverso durante la risposta di resistenza a ZYMV. Inoltre, la predizione dei geni R ha fornito ulteriori geni candidati potenzialmente implicati nella resistenza a ZYMV. La differente espressione genica, evidenziata in questo lavoro tra le due linee isogeniche, sarà utile per costruire un modello esplicativo sul processo di risposta dello zucchino a ZYMV. Infine, i risultati ottenuti saranno molto utili per velocizzare la selezione di cultivar locali di zucchino resistenti a ZYMV.
ABSTRACT

Cucurbits species are economically important crops grown worldwide. In many regions, the most serious threat to their production is Zucchini yellow mosaic virus (ZYMV). A monogenetic source of resistance to ZYMV was identified in *Cucurbita moschata* and introgressed in a *Cucurbita pepo*. However, in cultivated zucchini the major dominant gene indicated as *Zym-1* interact with other genes, making difficult the breeding transfer. Until now, not effective markers associated to ZYMV resistance have been found in *C. pepo*.

Main goal of this work was to investigate the mode of inherence of ZYMV resistance in zucchini and to identify molecular markers associated with resistance genes to ZYMV, in order to make easier, faster and economically sustainable the introgression of this resistance in zucchini local varieties ‘San Pasquale’. Moreover, a genome-wide transcriptional analysis of a resistant and a susceptible genotype to ZYMV was conducted in order to identify new candidate genes for resistance to ZYMV and to catalogue SNP polymorphic markers.

Two F₂ and three BC₁ segregating populations, obtained from the cross of ‘381e’ (resistant variety) with ‘True French’ and ‘San Pasquale’ (susceptible varieties) were assayed for ZYMV resistance. Phenotypic analysis highlighted a complex inheritance of resistance with the involvement at least of three genes (*Zym-1, Zym-2* and *Zym-3*). Molecular segregation analysis conducted using High Resolution Melt (HRM) technology on 3 SNP loci strongly suggests a relationship of these SNPs with resistance genes *Zym-1, Zym-2* and *Zym-3*. Moreover, nearly 3000 differentially expressed genes between ‘True French’ and ‘381e’ isogenic lines for resistance to ZYMV were identified in transcriptome of susceptible and resistant varieties inoculated with the virus. A high number of SNPs between the two varieties were also evidenced. A close co-localization of SNP1 with a CNL and an ATP-dependent RNA helicase, suggests that one of those genes could be a good candidate for exerting resistance function to ZYMV.

The study of ZYMV resistance response in zucchini at genetic and genomic level, performed in this work, can be useful for the genetic improvement of local variety as the susceptible ‘San Pasquale’ that have a good marketability in Campania region.
1. INTRODUCTION

1.1 Towards sustainable agriculture

Intensive agriculture practices have dramatically changed the ‘Earth’ ecosystem. In order to achieve maximum production with minimum efforts the modern agricultural system is based on high crop specialization, motorization and mechanization. The massive use of pesticides involves enormous damages to ecosystems leading to a gradual contamination of the soil and the water, the same that is used for irrigation, inducing resistance reactions by pests and destroying beneficial organisms. Therefore, a more sustainable agriculture system, to apply both in developed and in developing countries, is required to meet “the needs of the present without compromising the ability of future generations to meet their own needs”. Sustainable agriculture should maintains the adaptive capacity of farming systems (Hansen, 1996; Gold, 1994; Hansen and Jones, 1996; Park and Seaton, 1996). Agriculture is sustainable when it is ecologically and, economically viable, socially just, culturally appropriate, and based on a holistic scientific approach” (Madden et al., 1997). Although no one set of farming practices were established for sustainable agriculture, the cultivation of disease resistant varieties can meet the sustainable agriculture purpose (Horrrigan et al., 2002).

1.2 Cucurbita pepo

Morphologically, Cucurbita pepo is the most polymorphic species with respect to fruit characteristics (Duchesne, 1786; Naudin, 1856). The edible forms of this species can be grouped into eight morphotypes: Pumpkin, with round or nearly round fruits; Vegetable Marrow, with short, tapered and cylindrical-shaped fruits; Cocozelle, with long bulbous fruits; Zucchini, with uniformly cylindrical shaped fruits; Acorn, with furrowed, turbinate fruits; Scallop, with flat, scalloped fruits; Crookneck, with narrow, usually curved and warded, necked fruits and Straightneck, with short-necked or constricted fruits, usually warded (Paris, 1986). The great variability of types represented in the Botanical works of the 16th and 17th centuries, and the observations of the first European explorers in America, seem to indicate that Pumpkins, Scallops and Acorns were developed under the guidance of Native Americans in pre-Columbian times (Paris, 2001; Smith, 1997; Whitaker, 1947). The subsequent arrival in Europe of C. pepo fruits and seeds generated an extraordinary variability of new phenotypes through hybridization and recombination. New cultivars were developed, particularly the elongated forms of ssp. pepo: Vegetable Marrow, Zucchini and Cocozelle.
The great importance of the zucchini cultivation is due to the use of fruits for the greatly appreciated organoleptic characteristics that are a rich source of nutrients such as vitamins (vitamin A, C, folate) and minerals (potassium). About 24 million tons of zucchini (also reported as squash), together with pumpkins and gourds, were produced in the world in 2012 for a cultivated area of 1.8 million of hectares. China is the largest producer followed by India and Russian Federation. Italy, with around 520,000 tons, is the ninth producer country (FAOSTAT 2012; http://faostat.fao.org). In Italy, zucchini are among the first twenty commodities for production, covering more than 50% of European production, and is particularly developed in south regions (ISTAT 2007). The major production centres are located in Southern Italy (Sicily 2,920 ha; 1,338 Lazio ha; Puglia 1,510 ha; Calabria 1,342 ha; Campania 793 ha), but also in the North (Piemonte has 1,401, the Veneto has 1,027 and Emilia Romagna 116 ha) (ISTAT 2007).

1.3 Breeding for disease resistance in Cucurbita pepo

In the last century, breeders have made great efforts in zucchini breeding in order to develop new and improved cultivars and hybrids, nowadays among the main objectives of the genetic breeding there is also the recovery of local landraces. Breeding methods are based on crosses followed by selection pedigrees and backcross (Allard, 1960) and can be distinguish in pedigree and backcross methods. The first one, pedigree method, involves self-fertilization and selection for characters of interest in each generation and is commonly used when both of parental plants have positive characteristics. The backcross provides the selection of a parental showing a character of interest and then it is crossed with plants missing of that character and is commonly used to obtain plants resistant to several diseases. Disease resistance is considered the most important goal of summer squash breeding as C. pepo is highly susceptible to diseases (Whitaker and Robinson, 1986). Main zucchini diseases are caused by fungi and viruses as well as damages induced insects attack. Among fungi, powdery mildew, caused by Podosphaera xanthii or by the association of Erisyphe chioracearum and Sphaerotheca fuliginea, is a serious disease affecting the leaves, stems and fruits of cucumber and zucchini squash grown in greenhouses and in open field. On the other hand, insects, in particular Aphis gossypii, represents one of the major pests of cucurbits and ornamental plants inducing leaves deformation and stunted growth. In addition, insects represent a serious problem affecting Cucurbitaceae because they are responsible of the transmission of viruses, especially the Zucchini Yellow Mosaic Virus (ZYMV).

The potyvirus ZYMV is one of the most destructive and widespread viral pathogens on cucurbits worldwide (Zellning et al., 2014). It was first described in 1973 in northern Italy (Lisa et al., 1981).
ZYMV infection dramatically alters normal leaf and fruit shape, size and colour. Leaf symptoms include veinal chlorosis, vein banding, blistering, and strapleaf. Fruits display pronounced chlorotic bumps, malformations, and stunting. Mixed field infections of cucurbit species with ZYMV, PRSV, WMV and Cucumber mosaic virus (CMV) were reported to be common in cucurbit species and cause considerable damage worldwide and severe epidemics in cucurbit fields (Grafton-Cardwell et al., 1996; Lecoq et al., 1981). A study of ZYMV synergism was performed on the interaction of ZYMV with CMV, as these viruses are commonly found together in nature and can be transmitted by the same aphid (Lecoq et al., 1981) resulting in some cases in plants death (Wang et al., 2002).

A complete list of resistance genes involved in breeding programs has been identified in the Cucurbita species and published in 2005 by H. S. Paris. There are about 66 genes reported for C. pepo, 21 for C. moschata, 19 for C. maxima genes and 29 for the interspecific cross of C. maxima-C. ecuadorensis (25 out of these genes are enzymatic isoforms). One or two genes were also identified in 4 wild species (C. okeechobeensis, C. lundelliana, C. ecuadorensis and C. fetidissima HBK) and in several other interspecific crosses. Furthermore, a low number of resistance genes was introgressed in commercial zucchini cultivars from genotypes of related species, such as C. moschata Duchesne, and wild relatives, such as C. okeechobeensis (small) Bailey. These species represent an existing source of genetic variation to incorporate required traits, but their use is limited by cross-ability barriers, including reduced fertility and quasi-linkage.

1.3.1 Plant virus resistance mechanism

Plant viral diseases cause serious economic losses reducing yield and quality of crops and, although viruses are relatively simple genetic entities, much remain unknown about the mechanisms by which the many symptoms of disease are generated, and by which plants resist these effects.

The study of plant resistance raises many fundamental questions regarding the molecular, biochemical, cellular, and physiological mechanisms involved in the plant-virus interaction and the co-evolution. The first step in the study of genetics of viral resistance is to determine if the resistant response is inherited, and the number of genes involved. More than 80% viral resistance reported is monogenically controlled (the remaining shows oligogenic or polygenic control) (Kang et al., 2005). Only slightly more than half of all reported monogenic resistance traits show dominant inheritance. In most but not all cases, dominance has been reported as complete and dominant resistance is often associated with the hyper-sensitive response (HR) (Fraser, 1986).

The majority of characterized viral resistance genes from plants fall into the NBS-LRR class of resistance genes, such as HRT, RTM1, RTM2, RCY1, from Arabidopsis, and from solanaceous
hosts, N, Rx1, Rx2, Sw5, and Tm-22, all sharing structural similarity (Kang et al., 2005). However, the strong structural similarity does not ensure the same spectrum of action. A. thaliana RCY1 (resistance to C strain Y1) and HRT (HR to turnip crinkle virus) encode proteins that share 91% similarity (Takahashi et al., 2002), but confer resistance to unrelated viruses such as cucumber mosaic virus (CMV, a cucumovirus) and turnip crinkle virus (TCV, a carmovirus), respectively (Soosaar et al., 2005). In cultivated tomato, ToMV (Tomato mosaic virus) infections are controlled by the introgressed Tm-1, Tm-2 and Tm-22 genes. The Tm-22 resistance gene was shown to be durable (Hall et al., 1980; Pelham, 1966) and it has been cloned and well characterized (Lanfermeijer et al., 2003). Susceptible tomato plants transformed with the Tm-22 gene displayed resistance against ToMV infection and the resistance was conserved in all transgenic lines. In Citrus, it has been reported the resistant gene dominant gene Ctv-R that conferring broad-spectrum resistance against Citrus tristeza virus (CTV), one of the major pathogen of citrus (Rai et al., 2006). Seo and collaborators (2006) identified the TIR-NBS-LRR gene RT4-4 involved in a viral resistance response in common bean (Phaseolus vulgar cv. Othello). The functional analysis revealed that the RT4-4 gene in transgenic N. benthamiana lines is up-regulated in a non-virus-specific manner, although RT4-4 did not confer resistance to the reporter virus, it activated a resistance-like response (systemic necrosis) to Cucumber mosaic virus (CMV) (Seo et al., 2006). Studies on potyviruses resistance genes led to the identification of resistance genes corresponding to mutations in translation initiation factors, eukaryotic initiation factors 4E (eIF4E) and to a lesser extent, the eukaryotic initiation factor 4G (eIF4G) (Piron et al., 2010). The eIF4E gene mutation providing resistance to several Potyviridae family viruses has been identified in pepper (pvr1), pea (sbml), lettuce (mol (1), mol (2)), tomato (pot1), melon (nsv) and barley (rym4/5) (Robaglia et al., 2006; Ruffel et al., 2005). The modification in eIF4G is responsible for resistance of rice to Yellow mottle virus (Albar et al., 2006) and in Arabidopsis to Cucumber mosaic virus and Turnip crinkle virus (Yoshii et al., 2004).

Moreover, there are several examples of dominant and recessive genes that seem to control a wide range of viral genotypes. The most representative examples involve members of the Potyviridae. The I gene of common bean (Ali, 1950) was first reported to control a dominant resistance to ten different related potyviruses, Azuki mosaic virus, Bean common mosaic virus (BCMV), Bean necrotic mosaic virus, Blackeye cowpea mosaic virus, Cowpea aphid-borne mosaic virus, Passionfruit woodiness virus-K, Soybean mosaic virus (SMV), Thailand Passiflora virus, Watermelon mosaic virus (WMV), and Zucchini Yellow Mosaic Virus. As well in Cucumis melo a dominant gene (Ag) confer resistance to CMV transmitted by Aphis gossypii (Pitrat and Lecoq, 1980).
Genetic resistance is considered the most cost-effective and reliable approach to protect crops from virus infection. Several resistance genes have been tagged with molecular markers including RFLP, AFLP, RAPD, and various other PCR-based markers and these molecular approaches has been used for indirect selection via genotype, for localizing and isolating R genes in plant genomes.

1.4 Molecular marker in breeding for resistance to biotic stress

The construction of molecular maps and marker-assisted selection can aid the introgression of resistance to pathogens. Marker-assisted selection (MAS) provides opportunities for enhancing the response from selection because molecular markers can be applied at the seedling stage, with high precision and reductions in cost. The use markers associated to genes conferring monogenic resistances and quantitative trait loci (QTL) for quantitative disease resistances have been reported in several crop (St. Clair, 2010).

Traditional DNA-based methods include PCR-based methods and hybridization-based methods, i.e. restriction fragment length polymorphisms (RFLPs). PCR-based methods are extremely sensitive, often faster than other technologies, and are widely used in agriculture and food control (Grassi et al., 2006; Labra et al., 2004; Mane et al., 2006; Pasqualone et al., 2007). Random amplification polymorphic DNAs (RAPDs), amplified fragment length polymorphisms (AFLPs), inter simple sequence repeats (ISSRs), microsatellites or simple sequence repeats (SSRs), cleaved amplified polymorphic sequence (CAPS) and single nucleotide polymorphisms (SNPs). CAPS proves useful for genotyping, positional or map based cloning and molecular identification studies (Konieczny and Ausubel, 1993; Komori and Nitta, 2005; Spaniolas et al., 2006; Weiland and Yu, 2003).

Recently, SNP markers have become the most used markers in genetic characterization studies as well as in translational genomic (Varshney et al., 2014).

Single nucleotide polymorphisms are the most abundant forms of genetic variation among individuals of a species. SNPs are bi-allelic and allow an easy comparison between results collected from different laboratories using different chemicals and technologies. SNP markers were widely investigated in several crop. In corn, the frequency of polymorphism found in inbred lines is 1 SNP every 31 bp in the non-coding regions, and 1 SNP every 124 bp in the coding regions (Ching et al., 2002). In soybeans, genetic diversity studies in 22 different genotypes have detected the presence of 1.64 SNPs every 1000 bp in the coding regions and 4.85 SNPs every 1000 bp in the non-coding regions (Zhu et al., 2003). A research on barley from Kanazin et al. (2002) reports a frequency of 1 every 189 bp mutation. In Zucchini, Blanca and colleagues performed the first study based on SNP polymorphisms in 2011 identifying 9,043 high confidence SNPs between subsp. pepo cv. Zucchini
and subsp. *ovifera* cv Scallop, 4 SNPs per 100 bases in the coding regions. The popularity of SNPs as valuable and efficient molecular markers has increased the need to have more accurate SNPs detection methods. High resolution melting (HRM) analysis has been developed to detect SNPs in small PCR amplicons because it is an easy, require low-time (Hoffmann *et al*., 2007; Hung *et al*., 2008; Liew *et al*., 2004) and discriminates homozygous and heterozygous (Bennett *et al*., 2003; Herrmann *et al*., 2007; Stephens *et al*., 2008). The evolution of the HRM techniques from traditional melting curve analysis was implemented by the invention of new generation intercalating dyes (SYBR green I dye) and new instrumentation (Slinger *et al*., 2007). Melting curve analysis using fluorescent dyes has proven to be a highly sensitive method for mutation discovery and SNP genotyping (Liew *et al*., 2004; Cho *et al*., 2008; Smith *et al*., 2008).

The development of high-throughput methodologies allows the detection of many thousands of SNPs through genotyping platforms, facilitating the construction of high-density genetic maps (Gupta *et al*., 2008). Different SNPs genotyping technologies are now available: TaqMan® probes (Livak *et al*., 1995), FP-TDI (Chen *et al*., 1999), Invader (Olivier 2005), IPLEX (Wright *et al*., 2008), Kaspar (Nijman *et al*., 2008), SNApshot (Applied Biosystems; Sanchez *et al*., 2003), GoldenGate (Illumina, San Diego, USA; Fan *et al*., 2003) and Infinium (Illumina, San Diego, USA; Gunderson 2009). These genotyping techniques have been used extensively in humans (International HapMap Consortium, 2003), in several animal species (McKay *et al*., 2008; Kijas *et al*., 2008; Malhi *et al*., 2011) and are also available for plant species (Pavy *et al*., 2008; Hyten *et al*., 2008; Akhunov *et al*., 2009; Close *et al*., 2009; Muchero *et al*., 2009; Yan *et al*., 2009; Deulvot *et al*., 2010; Lepoittevin *et al*., 2010; Massman *et al*., 2011). The development of these new methods for the analysis of hundreds or thousands of SNPs at the same time allowed the possibility to build a dense SNP-genetic map in *Cucurbita pepo* (Esteras *et al*., 2012).

### 1.5 Sequencing and Next Generation Sequencing

In 1977, Maxam and Gilbert first reported an approach for DNA sequencing in which terminally labeled DNA fragments were subjected to base-specific chemical cleavage and the reaction products were separated by gel electrophoresis (Maxam and Gilbert, 1977).

In the last 30 years, the sequencing Sanger method, based on chain-terminating dideoxynucleotide analogs that caused base-specific termination of primed DNA synthesis (Sanger *et al*., 1977), had been the main approach to improve knowledge about genome sequences and, among the others, the conclusion of the first revision of the human genome in 2004 (International Human Genome Consortium, 2004).
From 2005, the use of massively parallel pyrosequencing platforms led the scientific community in the new era of high-throughput genomic analysis (Next Generation Sequencing-NGS). Starting about ten years ago, several instruments for sequencing were produced, such as Genome Sequencer from Roche 454 Life Sciences (www.454.com), the Solexa Genome Analyzer from Illumina (www.illumina.com), the SOLiD System from Applied Biosystems (www.appliedbiosystems.com), the Heliscope from Helicos (www.helicos.com) and the commercialized Polonator (www.polonator.org). All these machines are able to give higher yield with simpler process and to produce sequences in a cost-efficient way. Although each instrument is different from the others, all massively parallel sequences machines share, more or less, the same way of action (Mardis, 2011).

First, the amplification of the library fragments by PCR, involving multiple primer pairs. Following the PCR, the resulting fragments have platform-specific adapters ligated to their ends to form a library that is ready for sequencing. After sequencing, the followed step is the computationally conversion of image data into sequence reads, the so-called “base calling”, to generate read sequences and to highlight the error probability–related quality scores for each called base. The quality values calculated during the sequencing is related to the historically relevant Phred score, introduced in 1998 for Sanger sequence data (Ewing et al., 1998a; Ewing et al., 1998b). The Phred score quality value, $q$, uses a mathematical scale to convert the estimated probability of an incorrect call, $e$, to a log scale: $q=-10 \times \log_{10}(e)$. Miscall probabilities of 0.1 (10%), 0.01 (1%), and 0.001 (0.1%) yield Phred scores of 10, 20, and 30, respectively. This score resulting to be crucial for eliminating low quality reads and bases, improving alignment accuracy and determining consensus-sequence and variant calls (Li et al., 2008).

Finally, the obtained reads are used to make an assembly, or a hierarchical data structure, that maps the sequence data in order to reach a putative reconstruction of the target. Reads are grouped in contigs (provide a multiple sequence alignment of reads plus the consensus sequence) and contigs into scaffolds (called also supercontigs or metacontigs, define the contig order and orientation and the sizes of the gaps between contigs).

To date, several genomes are sequenced de novo or resequenced (Wheeler et al., 2008; Margulies et al., 2005; Pearson et al., 2007; Smith et al., 2008; Quinn et al., 2008; Satkoski et al., 2008; Borneman et al., 2008; Wang et al., 2008) and, moreover, NGS platforms have been used to detect new structural variation, such as deletions, insertions and rearrangements (Korbel et al., 2007; Campbell et al., 2008; Kim et al., 2008; Chen et al., 2008).
1.5.1 RNA-Seq

One of the most popular NGS approaches is the RNA sequencing (RNA-seq) employed to map and quantify transcripts in biological samples. This technology include the analysis of a population of RNA where each molecule, with or without amplification, is sequenced in a high-throughput manner to obtain short sequences from one end (single-end sequencing) or both ends (pair-end sequencing). The alignment of reads is the consequential step, if a reference genome is available, or *de novo* assembling of transcriptome, a transcription map that consists of both the transcriptional structure and level of expression for each gene is required (Wang *et al*., 2009).

By counting reads that map on sequenced exons (assembled in transcripts corresponding to genes) is possible to obtain the expression levels of each transcripts. The quantity of mapped reads per transcript must be normalized in order to compare the expression levels of transcripts within the same sample and between different samples. The most used normalization methods are RPKM/FPKM (reads or paired-end fragments per kilobase of exon model per million mapped reads) (Mortazavi *et al*., 2008), TMM (trimmed mean of M-values) (Robinson *et al*., 2010) and CPM (count per million) (Robinson *et al*., 2010).

The biggest revolution mediated by the sequencing of RNA is the ability to give an accurate view about gene expression levels and, on the other side, the possibility to detect and study the transcriptome for non-model organisms that have no genomic sequences determined (Vera *et al*., 2008; Marioni *et al*., 2008). Moreover, the RNA-seq strategy can also reveal sequence variations, such as SNPs, isoform and novel splice junctions, and information of antisense regulation and intragenic expression (Cloonan *et al*., 2008; Morin *et al*., 2008; Carninci *et al*., 2005; Nagalakshmi *et al*., 2008; Graveley *et al*., 2011; Trapnell *et al*., 2010).

Although the RNA sequencing is a relatively new approach, it has been already applied to several crop plants (Bagnaresi *et al*., 2012; Gao *et al*., 2013; Mesarich *et al*., 2014).
1.6 Aims of the work

Main goal of this work was to investigate the mode of inherence of ZYMV resistance in zucchini and to identify molecular markers associated with resistance genes to ZYMV, in order to make easier, faster and economically sustainable the introgression of this resistance in local varieties of zucchini, as ‘San Pasquale’. Second aim of our work was to conduct a genome-wide transcriptional analysis of a resistant and a susceptible genotype to ZYMV, in order to identify new candidate genes for resistance to ZYMV, to perform a R genes prediction and to catalogue new SNP markers linked to the genes conferring ZYMV resistance. Results obtained constitute an important compendium of methods and information that can improve selection procedures employed in *Cucurbita pepo* breeding program.
2. MATERIALS AND METHODS

2.1 Plant materials

Four zucchini varieties (Nano Verde di Milano, True French, 381e and 968Rb), four Cocozzelle varieties (Romanesco, Ortolana Di Faenza, San Pasquale and Bianca di Trieste) and one Pumpkin accession (Tondo Chiaro di Nizza) were used for genotyping assay. The susceptible open-pollinated zucchini ‘True French’ (TFR) (Thompson & Morgan, Ipswich, England, 1978) was crossed with its near-isogenic resistant counterpart, Accession ‘381e’ indicated from hereafter as ‘381e’ to obtain a F2 populations. The latter variety had been derived from six generations of backcrossing to ‘True French’, selecting for resistance, followed by four successive generations of self-pollination and selection for resistance (Paris and Cohen, 2000). ‘San Pasquale’ referred hereafter as ‘SPQ’ obtained from La Semiorto Sementi, Sarno, Italy, is an open-pollinated cocozelle cultivar having extremely long, striped fruits and susceptible to ZYMV, was also crossed with ‘381e’ to produce F2 populations. Three BC1-F2 populations were obtained from the cross of different F2 ‘SPQx381e’ individuals (plants 64, 47 and 28) with the recurrent parent ‘SPQ’.

2.2 Resistance to ZYMV in vivo assay in segregating populations

Seeds of parental and filial generations were sown in multi-cellular trays consisting of 4 cm diameter pots that had been filled with peat, one per pot. After the seedlings were transferred into pots 15 cm diameter and were grown under glasshouse condition at 22-24°C using supplement lighting to maintain 12 photoperiod at the Research Centre for Plant Pathology (CRA-PAV) in Rome. Each individual plant was assigned an identification number.

An isolate of ZYMV from a naturally infected plant of field-grown summer squash was used for experiments. This isolate cause the typical symptoms of ZYMV disease including yellow mosaic, vein banding, blistering and malformation of leaves. Symptomatic leaves of artificially infected zucchini plants were crushed, and the raw juice was extracted at a ratio 1:10 w/v in 0.1 M phosphate buffer pH 7.2. Both cotyledons of test plants, sprinkled with the abrasive powder “celite”, were inoculated with approximately 20 μl of the diluted extract, and subsequently washed with distilled water. Symptoms were observed and recorded from 6 to 30 days after inoculation (dpi), when the plants had developed 6–7 true leaves. Plants each of the three parents as well as F1 plants were used as resistant, susceptible and heterozygous controls. According to the symptoms observed after 12 days after inoculation and based on what reported by Paris and Cohen (2000), the plants were classified in three groups. Group A, asymptomatic plants or with few yellow spots on new leaf.
Group B, plants with yellow spots spread on new entire leaf and with symptoms of "vein banding" after 12 dpi and then recovery after 25 dpi. Group C, plants with symptoms of "vein banding" and mottle on leaves or clearly susceptible plants with leaf malformation.

To ensure that the asymptomatic plants were really resistant and not simply escaped by inoculation of the virus, a serological ELISA test was performed. The presence of the virus into infected plants would prove by an intense serological reaction (yellow) compared to healthy controls.

2.3 Molecular analysis

2.3.1 DNA isolation
In the young leaves, using the protocol described by Fulton et al. (1995), was isolated total genomic DNA of parental and segregating population samples. Each sample was represented by a bulk of three leaves. The DNA amount was quantified spectrophotometer by NanoDrop ND-1000 Spectrophotometer (NanoDrop Technologies) and the quality was checked electroforetically with a 1.2% agarose gel staining with a SYBR® Safe DNA Gel Stain (Takara). Samples were then diluted to a 15-25 ng/μl working concentration and stored at -20°C.

2.3.2 Genotyping assay
An Illumina GoldenGate genotyping assay, based on a 384 SNPs panel, was performed on nine varieties of Cucurbita pepo sbp pepo. These varieties include representatives of three morphotypes of the subspecies pepo such as Zucchini varieties (Nano Verde di Milano and three isogenic lines True French, 381e and 968Rb); Cocozzelle varieties (Romanesco, Ortolana Di Faenza, San Pasquale and Bianca di Trieste) and Pumpkin accession (Tondo Chiaro di Nizza). The genotyping was performed through a Principal Component Analysis (PCA) obtained using GenAlEx 6.41 and MEGA5 software.

2.3.3 CAPS analysis
The F2 segregating population obtained from ‘True French’ and ‘381e’ was tested with a CAPS marker designed thanks to the GoldenGate platform data. The CAPS analysis consisted of a PCR reaction and subsequent digestion with BglII restriction enzyme that recognizes A^GATCT sites. PCR amplification was carried out in a final volume of 25 μL, Buffer 1x, 0.1 mM dNTPs, 0.2 μM, Taq polymerase 0.5 U, and 0.2 μM each primer, DNA 25-35 ng. Amplification was performed
under the following program: 30 s at 95° C; 40 cycles of 10 s at 95° C, 30 s at 57° C, and 30 s at 72° C; and extra extension for 1 min and 30 s at 72° C. 15 µL of amplified products was treated for 3 hours at 37° C with 1 U BglII restriction enzyme in a final volume of 25 µL. Polymorphism was detected by separating the whole volume of treated DNA on 2% agarose gel (1x Tris-Acetate EDTA) containing SYBR® Safe DNA Gel Stain (Takara), and visualizing it under a UV transilluminator. We used 1Kb+ (Invitrogen) fragments as size markers.

2.3.4 HRM analysis

Real-Time PCR mixture was prepared using 15-25 ng of DNA, 0.6 U of Takara ex Taq™ (Takara), 20 mM of Mg2+, 2.5 µl of 10X ex Taq Buffer 10 mM, 2.5 mM dNTPS and primers were mixed to the final concentration of 10 uM. The intercalating dye used was the EvaGreen (Biothium) at final concentration of 0.5X. Real-Time PCR reactions and HRM analysis were performed on Rotor-Gene 6000™ (CorbettResearch, Cybeles, Thailand). The Real-Time PCR profile comprised of one initial cycle of 30 s at 95°C and followed by 40 cycles of 10 s at 95°C, 30 s at 58-60°C, 30 s at 72°C. After Real-Time PCR amplification, HRM was performed using melting profile from 70° to 90°C rising at 0.1°C per sec. The software provided by the company normalized the melting curves. The specificity of primers used in the experiment was confirmed by a single amplicon melting analysis. Data from HRM analysis was determined using Rotor-Gene 6000 series 1.7 software. All of the analyzed data was achieved according to the manufacturer’s protocol (HRM guide www.appliedbiosystems.com).

2.4 Statistical analysis goodness of fit

ZYMV resistance phenotypic segregation of F2 ‘TFRx381e’ and F2 ‘SPQx381e’ population was analyzed by chi-square test to establish the degree of coincidence between the expected and the obtained segregation ratio 45:19. The results from HRM analysis for all molecular markers were subjected to 3:1 χ2 analysis, too. Moreover, the segregation into resistant and susceptible genotypes was also calculated as the frequency of resistant plants f(R) – number of resistant plants/number of tested plants per combination. In the two BC1-F2 populations the expected ratio of resistant/susceptible was 1:1 both in phenotypic and genotypic evaluations. Furthermore, a Mantel test (Mantel, 1967) was executed between phenotypic and genotypic data to find a correlation between the two trends. In order to obtain the phenotypic and genotypic matrices was performed the Ascendant Hierarchical Coefficient (HAC) by the XLStat tool. MEGA5 tool was
used to perform, on the obtained matrices, the dendrograms based on the Euclidean distance between the distributions. The Pearson’s correlation coefficient was evaluated on the two matrices through a correlation/association test (Mantel test).

2.5 Analysis of genome scaffolds containing ZYMV markers

In order to identify genes physically closer to our markers, a functional prediction of protein comprising domains involved in the defense mechanisms was performed. Firstly, the three unigenes extracted from the GoldenGate panel have been searched on the genome and obtaining the scaffolds containing the unigenes. GenScan tool was used to translate the scaffolds sequences in the peptides in order to verify, via HMM profiling using InterProScan software version 4.8 (Zdobnov and Apweiler, 2001), the presence of conserved domains and motifs characteristic of proteins (NBS; LRR; TIR; KINASE; SERINE/THREONINE). In this analysis, recovered sequences were compared with the following databases: Hidden Markov Model Panther (HMPanther), Hidden Markov Model Tigr (HMMTigr), patternScan, FPrintScan, HMMPIR, ProfileScan, High-Quality Automated and Manual Annotation of Microbial Proteomes (HAMAP), SignalPHMM PROSITE, SuperFamily PRINTS (Fingerprint database), HMMPfam (Protein family; http://pfam.wustl.edu/hmmsearch.shtml), Blast Protein Domain database (BlastProDom), and Hidden Markov Model Simple Modular Architecture Research Tool (HMMSMART) protein motif analyses (http://smart.embl-heidelberg.de/; Schultz et al., 1998). The TMHMM database was also accessed to verify the presence of transmembrane regions of the candidate proteins belonging to the class of transmembrane receptor (RLK and RLP proteins). The coils program (Lupas et al., 1991; http://www.ch.embnet.org/software/COILS_form.html) was used to detect CC domains in CNL proteins using a threshold = 0.9 followed by visual inspection.

InterPro is a resource that provides functional analysis of protein sequences by classifying them into families and predicting the presence of domains and important sites. To classify proteins in this way, InterPro uses predictive models, known as signatures, provided by several different databases (referred to as member databases) that make up the InterPro consortium. The software package InterProScan allows sequences to be scanned against InterPro's signatures (more information are available in http://www.ebi.ac.uk/InterProScan/).
2.6 Transcriptomic analysis

2.6.1 RNA isolation, quantity and quality evaluation

The sampling was conducted on leaves of 3 groups of 6 plants each at 12th and 22th days after inoculation (dpi) on the two susceptible and resistant genotypes. Collected samples were immediately submerged in liquid nitrogen and then stored at -80°C. The total RNA extraction was performed on 100 mg of frozen zucchini leaves using TRIZOL (Invitrogen) protocol (Chomczynski and Mackey, 1995). A treatment with DNase I to remove DNA contaminations was done on total RNA. RNA samples were valuated quantitatively and qualitatively by NanoDrop ND-1000 Spectrophotometer (Nano-Drop Technologies, Wilmington, DE, USA) and by Bioanalyzer (Agilent Technologies). RNA integrity was checked by horizontal electrophoresis on a 1.2% (w/v) agarose gel prepared without addition of ethidium bromide. Two μg of each sample were prepared with 20 μl of 10 X RNA Loading Buffer composed by 400 μl Formamide, 10 μl 37% formaldeyde, μl loading buffer 10X (50% glycerol, 0.25% w/v bromophenol blue, 0.25% w/v xylene cyanol; Sigma), and 1. μl of 10 mg/μl ethidium bromide and than denatured at 6 °C for 15 min. Gel visualization was performed using UV light (UV Gel Doc BIORAD). In order to obtain high quantity of RNA for each samples, total RNA from the six replicate was pooled for each 12 samples.

The samples, which have met the required standards of quality, were subjected to sequencing at the LabMedMolGe (Laboratory of Molecular Medicine and Genomics Department of Medicine and Surgery, University of Salerno).

2.6.2 RNA- Sequencing

Total purified RNA was converted to cDNA libraries and sequenced on Illumina HiSeq1500 platform at the LabMedMolGe (Laboratory of Molecular Medicine and Genomics Department of Medicine and Surgery, University of Salerno). The process generates millions (36 M) of short (100 bp) reads sequenced from both ends of each cDNA fragment (paired-end sequencing). The raw data for each sample consist of a long list of short sequences with associated quality scores (fastq format) (Ewing et al., 1998a; Ewing et al., 1998b). To evaluate the quality of the sequences the FastQC software was used (http://www.bioinformatics.babraham.ac.uk/projects/fastqc).
2.6.3 Functional annotation of reference transcriptome

Transcriptome (version 3.0), was mapped to the genome (version 2.0), in order to obtain the coordinates of the transcripts/genes over the genome. The PASA tool (version r20140417), which calls BLAT in order to align the transcripts over the genome and get the coordinates of the best hits (http://pasa.sourceforge.net), was employed. To perform the functional annotation of the reference transcriptome the BLAST2GO tool was used at the default parameters (Conesa et al., 2005).

2.6.4 RNA-Sequencing data analysis

RNA-seq analysis was performed in three different steps: Reads processing; Statistical analysis of the data; Functional annotation of the identified loci. The high quality reads (obtained after trimming) were aligned against the *Cucurbita pepo* transcriptome (version v3) using TopHat v.2.0.13 (http://ccb.jhu.edu/software/tophat). Bowtie2 (included in TopHat) was selected as read aligner due to its high sensitivity. High quality reads were aligned against the transcriptome using minimum insert-length as zero and maximum insert-length as 500 (or more). The resulting alignment files (.bam files) were filtered for quality (quality score > 30) and duplicates. The obtained files were used as input for the calculation gene expression values. The output file includes one column for the transcript ID and one column for expected-read-counts for each sample. Data counts were than normalized using the RPKM method (Mortazavi et al., 2008). All the statistical analyses were performed with R (version 3.1.0).

The overall quality of the experiment was evaluated, on the basis of the similarity between replicates, by a PCA analysis. Using this approach, we detected if any of the samples does not behave like the other replicates (outliers). The similarity between replicates was evaluated as well by the calculation of Euclidean distance between the samples and hierarchical clustering. The clustering of the samples was repeated after the removal. Once the consistency of biological replicates were evaluated, and divergent samples (outliers) were discarded, differential expression analysis was performed. The final dataset was composed by all the samples passing the Quality Control step.

The identification of the differentially expressed genes was performed with the package EBSeq (version 1.5.3). Sample 381-12dpi versus the sample TF-12dpi, 381-22dpi versus 381-12dpi, 381-22dpi versus TF-22dpi, TF-22dpi versus TF-12dpi have been compared. In addition, a more general analysis has been performed in order to identify DE transcripts in ‘381e’ genotype respect to ‘TFR’ genotype. In this experiment, the time was not treated as factor. The transcripts were scored at q-value less than 0.05. A further filter was applied by selecting those with a fold change higher than
1.5-fold (UP-regulated) and lower than -1.5-fold (DOWN-regulated). The attachment of biological information to identified differential expressed transcripts was performed using the functional annotation file of the *Cucurbita pepo* transcriptome (version 3.0).

In order to detect new SNP markers on the transcripts of the susceptible ‘TFR’ and resistant ‘381e’ accessions, a SNP calling analysis against the *Cucurbita pepo* reference genome was performed. SNP calling, based on the two filtered .bam files, produced two VCF (Variant Call Format) files (http://www.broadinstitute.org/igv/viewing_vcf_files) with the SNPs of each genotype in comparison to the genome. To obtain the SNPs between both genotypes we have found the common and unique SNPs between both VCF files, removing at the same time SNPs at low quality/coverage and inconsistent SNPs.

The protein sequences of our genes were interrogated by MATRIX RELOADED (Matrix-R) tools that, via HMM profiling, automatically retrieves, annotate and classified plant resistance genes. The protein sequences of the genes identified, were examined in order to underscore typical R gene domains. The output was a list of transcripts with a hit to any resistance domain Matrix-R was launched on transcripts with an ORF > 60 amino acids.
3. RESULTS

3.1 Genetic analysis of resistance to ZYMV

3.1.1 Phenotypic evaluation of resistance in segregating populations

Two F₂ populations deriving from the cross ‘True French’ x ‘381e’ (F₂ TRF) and ‘San Pasquale’ x ‘381e’ (F₂ SPQ) and three backcross populations obtained crossing F₂ ‘SPQ’ progenies with susceptible variety ‘SPQ’ were screened for resistance to ZYMV. Inoculated plants scored from 6 to 30 days post inoculation (dpi) exhibited wide scattered reaction to virus. Accession ‘381e’ and a number of resistant segregating progenies showed chlorotic spots, mild mottling then recovering or only a few chlorotic dots on new leaves. ‘True French’, ‘San Pasquale’ and all susceptible segregating plants showed severe mosaic, blistering and distortion on leaves and reduced growth.

As regards of ‘TFRx381e’ F₂ progeny, consisted of 175 individuals, 122 plants (70%) were classified as resistant, 102 plants showed the same symptoms of resistant parent, with mild mosaic and yellow spots at initial development stage (1-3rd leaf) and subsequent development of asymptomatic leaves (Group B). While, 20 plants showed no macroscopic symptoms on all leaves observed and classified as group A. Frequency of resistant plants on total of plants observed was calculated as f (R) = 0.70. The symptoms of the susceptible parent ‘TRF’ with malformations, mosaic and reduction of development were visualized in 53 individuals, about 30% of the full population screened. These individuals, exhibiting growth reduction, severe mosaic, blistering and leaf distortion were pooled in group C as previously described in materials and methods section. The observed segregation fitted with both the 3:1 and 45:19 ratio (Table 3.1), which suggested that one major gene control the resistant reaction in ‘381e’ but its action could be modulated from at least from other two other genes.

Also in the F₂ population ‘San Pasquale’ x ‘381e’ (n = 96), the segregation ratio was in accordance with a genetic control mediated from a major gene, putatively modulated from other genes. Phenotypic analysis of this population showed the occurrence of 62 resistant plants and of 34 susceptible, f (R) = 0.65. In particular, out of 62 plants judged resistant, 17 belong to the group A and 45 to the group B.

The F₂ individuals used to produce the backcross populations were carefully selected according to their allelic composition at putative loci Zym-1 (A), Zym-2 (B) and Zym-3 (C) revealed previously by the molecular analysis carried out on the ‘SPQx381e’ F₂ population. The genotype ‘64-F₂’ was characterized by a heterozygous locus Aa and two dominant homozygous loci BB and CC,
genotype ‘47-F2’ had an allelic configuration Aa, bb, cc, ‘28-F2’ is a triple heterozygote (Aa, Bb, Cc).


The results obtained confirm the previsions in the ‘47-F2’ and ‘28-F2’ backcross. In particular, BC1-F2 ‘47-F2xSPQ’ population evidenced 23 susceptible plants and 29 with symptoms of tolerance, on a total of 52 plants tested, f (R) = 0.56. In BC1-F2 ‘28-F2xSPQ’ out of 68 tested plants, 36 showed signs of susceptibility, 32 symptoms of tolerance, f (R) was 0.47. Finally, out of 68 backcross plants obtained from ‘64-F2’, 49 (72% instead of 50%) showed symptoms of susceptibility, 19 showed tolerance (28% instead of 50%), in the juvenile stages (20 dpi). Frequency of resistant plants was 0.28. Resistant plants were kept for further crosses useful to transfer the ZYMV resistance in the ‘San Pasquale’ genotype.

3.1.2 Molecular screening of parental lines

A 384 Illumina GoldenGate platform designed on a Zucchini Unigene-Dataset (Blanca et al., 2011) was used in order to analyze the genetic relationships of three parental lines and of six lines used as control. The GoldenGate genotyping assay allowed us to identify 39% of polymorphic SNPs among varieties tested. Based on SNPs profiles obtained a PCA profile was build up. The phylogenic distance and the correlation between the nine analyzed cultivars was evaluated as showed in figure 3.1. Only six SNPs detected variation between the two isogenic lines, ‘381e’ (resistant to ZYMV) and ‘True French’ (susceptible to ZYMV). Out of six markers used to develop a Real-Time PCR-HRM assays, three produced a clear melting curve.
Figure 3.1: Phylogenetic relationships between nine *Cucurbita pepo* cultivars. In the red circle on the left were pooled the three isogenic lines ‘True French’, ‘968Rb’ (susceptible) and ‘381e’ (resistant) and on the right the cultivar ‘San Pasquale’ belonging a group of cocozelle.

3.1.3 Genotyping of segregating populations

HRM assay with SNP1, SNP2 and SNP3 carried out on 5 segregating populations, characterized for resistance to ZYMV allowed a quick and efficient discrimination between individuals homozygous resistant, susceptible homozygous and heterozygous resistant at loci tested.

Thanks to the molecular analysis, we evaluated the putative association of SNP markers to three genes (*Zym-1, Zym-2 and Zym-3*) involved in resistance to ZYMV in two F2 populations and three backcross F2 populations.

The subset of three SNPs were investigated on 174 F2 ‘TFRx381e’ individuals, 91 F2 ‘SPQx381e’ individuals, 60 BC1-F2 ‘64-F2xSPQ’, 49 ‘47-F2xSPQ’ and 65 of ‘28-F2xSPQ’ individuals. The assay performed on the five segregating populations allowed the clear discrimination of allelic state at three loci in all individuals tested. High-Resolution Melting analysis revealed significant differences in melting curve shape that correlate SNP to the presence of a given allele. Therefore based on this assay it was possible to discriminate dominant homozygotes (AA), heterozygotes (Aa) and homozygous recessive (aa) individuals for each locus. Representative normalized melt curves of SNP1 marker are shown in figure 3.2.
Figure 3.2: High Resolution Melting Real-Time PCR. HRM profiles of SNP1 marker in a segregating population of zucchini plants. Normalized melting plot between resistant homozygous R/R, susceptible homozygous r/r and resistant heterozygous R/r are shown. Representative profiles of the melting curves of amplicons for F2 population.

HRM analysis performed on the ‘F2 TFR’ population revealed a segregating ratio of 130 A- plant to 44 aa plants, f (A-) = 0.75 and on the F2 obtained from the ‘F2 SPQ’ showed a segregating ratio of 66 A- to 25 aa samples, f (A-) = 0.73. In these F2 populations, the resistant individuals were characterized by the configurations A-B-C-, AaBBcc, A-bbC-, A-bbcc, and the susceptible were aaB-C-, aaB-cc, aabbC-, aabbcc. These 27 allelic configurations are summarized in figure 3.3.

In the BC1-F2 ‘64-F2xSPQ’ the f (A-) calculated was 0.47. On 60 analyzed plants, 28 resulted with the allelic configuration AaBbCc and 32 aabbcc. While in the ‘47-F2xSPQ’ backcross the resistant plants was 21 (Aabbcc) and the susceptible 28 (aabbcc) with an f (A-) = 0.43. Lastly, in the BC1-F2 ‘28-F2xSPQ’ there was 36 resistant subdivided in ¼ of individuals AaBbCc, ¼ of individuals AaBbCc, ¼ of individuals AaBBcc and ¼ of individuals AaBBcc as well as the 29 susceptible individuals were subdivided in ¼ aaBbCc, ¼ aaBbCc, ¼ aabbCc and ¼ aabbcc for a total of 65 (f (A-) = 0.55). The results obtained by HRM analysis for each marker were subject to a chi-square test for a single dominant gene segregation ratio 3:1 and for three dominant genes segregation ratio 45:19. This test has highlighted a strong correlation between SNP observed frequency and expected frequency for both ratio.
Figure 3.3: F2 expected allelic configuration of three complementary genes involved in resistance mechanism. In green and red are labeled the resistant and susceptible genotypes, respectively. In this work, the genotypes Aabbcc* and Aabbcc* showed resistance variations among individuals.

The SNP1 marker showed a strong correlation with phenotypic data and it was used to develop also a CAPS marker able to validate results obtained by HRM assay on 167 F2 ‘TFRx381e’ individuals.

The PCR reaction produced amplicons size of approximately 400bp. The BglII restriction enzyme digestion allowed us to discriminate susceptible from resistant samples. In particular, the susceptible samples have produced two fragments of 200bp (displayed on an agarose gel as a single band), those judged resistant a 400bp fragment, whereas the heterozygous samples produce both fragments (Figure 3.4).

Figure 3.4: A Cleaved Amplified Polymorphism Sequence (CAPS) marker. The polymorphism of the marker SNP1 is shown in segregating population F2 generated crossing ‘381e’ (resistant parent) and ‘True French’ (susceptible parent). In lane 5: resistant homozygous genotypes (R/R 400 bp); in lane 1, 3, 9, 10, 11 and 12: susceptible homozygous genotype (r/r – 200 bp); in lane 2, 4, 6, 7 and 8: resistant heterozygous genotypes (R/r - 400 bp and 200 bp), M: ladder (1Kb*, Invitrogen).
3.1.4 Statistical analysis goodness of fit

‘True French’ x ‘381e’ F₂ population phenotypic evaluation showed 122 resistant samples and 53 susceptible samples, in accordance with the expected 45:19 ratio ($\chi^2 = 0.03$ and $P = 0.86$). The f (R) number of resistant plants/total number of tested plants is 0.70 as shown in table 3.1. In addition, the ratio of 62 resistant on 34 susceptible individual observed in the cross between ‘San Pasquale’ and ‘381e’ statistically fitted with an expected ratio 45:19 ($\chi^2 = 1.51$ and $P = 0.22$). In this two F₂ population, the observed ratio also fitted with the 3:1 segregating expected ratio, but with lower value at $P > 0.05$ (Table 3.1). The 45:19 ratio was tested assuming that three complementary dominant genes are involved in resistance to ZYMV (Paris and Cohen, 2000).

The phenotypic evaluation of three backcross populations showed an observed ratio of 19 resistant and 49 susceptible in the ‘64-F₂’ derived backcross population, 29:23 in the ‘47-F₂xSPQ’ and 32:36 in the ‘28-F₂’. In the latter two, the observed ratio fitted with the 1:1 expected ratio.

Phenotypic evaluations suggest that the response to ZYMV in a first stage of resistance process require the participation of three genes. In particular, one major gene Zym-1 and two minor genes Zym-2 and Zym-3, which may have a complementary effect, producing an F₂ ratio of 45:19 and a BC₁-F₂ ratio of 1:1 seems to be involved. Analyzed plants were considered resistant assuming that the resistant genotypes should have at least one allele of the locus Zym-1 dominant (Zym-1/), complementary genes Zym-2 and Zym-3 (Zym-2/ and/or Zym-3/) could be necessary to keep resistance in later stage of development.

<table>
<thead>
<tr>
<th>ID_Population</th>
<th>No. pt Resistant</th>
<th>No. pt Susceptible</th>
<th>No. pt Total</th>
<th>Expected ratio (R:S)</th>
<th>$\chi^2$ (P)</th>
<th>f (R)</th>
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<tr>
<td>F₂ (TFRx381)</td>
<td>122</td>
<td>53</td>
<td>175</td>
<td>45:19</td>
<td>0.86</td>
<td>0.70</td>
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<tr>
<td>F₂ (SPQx381)</td>
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<td>34</td>
<td>96</td>
<td>45:19</td>
<td>0.22</td>
<td>0.65</td>
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<tr>
<td>F₂ (TFRx381)</td>
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<td>175</td>
<td>3:1</td>
<td>0.11</td>
<td>0.70</td>
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<tr>
<td>F₂ (SPQx381)</td>
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<td>34</td>
<td>96</td>
<td>3:1</td>
<td>0.02</td>
<td>0.65</td>
</tr>
<tr>
<td>BC₁-F₂ (64-F₂xSPQ)</td>
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<td>68</td>
<td>1:1</td>
<td>0.001</td>
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</tr>
<tr>
<td>BC₁-F₂ (47-F₂xSPQ)</td>
<td>29</td>
<td>23</td>
<td>52</td>
<td>1:1</td>
<td>0.71</td>
<td>0.56</td>
</tr>
<tr>
<td>BC₁-F₂ (28-F₂xSPQ)</td>
<td>32</td>
<td>36</td>
<td>68</td>
<td>1:1</td>
<td>0.88</td>
<td>0.47</td>
</tr>
</tbody>
</table>
Molecular results obtained with the HRM analysis conducted were also subject to chi-square test. For each marker was calculated the segregation ratio for a single dominant gene. Assuming that also the samples carrying the genotype Zym-1/Zym-1 zym-2/zym-2 zym-3/zym-3 and Zym-1/zym-1 zym-2/zym-2 zym-3/zym-3 were resistant, the segregating expected ratios were 3:1 in the two F2 populations and 1:1 in the BC1-F2 populations. The chi-square test performed on dataset obtained by the molecular analysis confirmed this expected ratios in all five cases suggesting a close linkage between analyzed SNPs and ZYMV resistance genes. Moreover, in order to envisage the tendency of genetic markers to co-segregate with the phenotypic data was calculated the percent of co-segregation between the genotype and the phenotype. Co-segregation was ascertained in all five population with a minimum of 47% in the BC1-F2 ‘47-F2xSPQ’, considering as susceptible the genotype Zym-1/Zym-1 zym-2/zym-2 zym-3/zym-3 and Zym-1/zym-1 zym-2/zym-2 zym-3/zym-3, to a maximum of 89% in the ‘TFRx381e’ F2. Table 3.2 reports the results of a chi-square goodness of fit test and the percent of co-segregation calculated between the genotype and the phenotype. A ratio of 3:1 segregation was observed for each F2 segregating populations reflecting a close linkage between analyzed SNPs and ZYMV resistance genes.

Table 3.2: Based on the genotypic data, chi-square test was calculated for 45:19 and 3:1 expected ratio in the F2 populations and for 1:1 expected ratio in BC1-F2 populations. The ratio labeled with * report the calculated $\chi^2$ considering as resistant the genotype Zym-1/Zym-1 zym-2/zym-2 zym-3/zym-3 and Zym-1/zym-1 zym-2/zym-2 zym-3/zym-3. The f (A-) and the percent of co-segregation of genotype vs phenotype was calculated, too.

<table>
<thead>
<tr>
<th>ID_Population</th>
<th>No. pt</th>
<th>No. pt</th>
<th>No. pt</th>
<th>Expected</th>
<th>Co-segregation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Resistant</td>
<td>Susceptible</td>
<td>Total</td>
<td>ratio (R:S)</td>
<td>genotype vs phenotype (%)</td>
</tr>
<tr>
<td>F2 (TFRx381)</td>
<td>130</td>
<td>44</td>
<td>174</td>
<td>3:1*</td>
<td>0.93</td>
</tr>
<tr>
<td>F2 (SPQx381)</td>
<td>66</td>
<td>25</td>
<td>91</td>
<td>3:1*</td>
<td>0.59</td>
</tr>
<tr>
<td>F2 (TFRx381)</td>
<td>130</td>
<td>44</td>
<td>174</td>
<td>45:19*</td>
<td>0.20</td>
</tr>
<tr>
<td>F2 (SPQx381)</td>
<td>66</td>
<td>25</td>
<td>91</td>
<td>45:19*</td>
<td>0.64</td>
</tr>
<tr>
<td>F2 (TFRx381)</td>
<td>121</td>
<td>53</td>
<td>174</td>
<td>3:1</td>
<td>0.10</td>
</tr>
<tr>
<td>F2 (SPQx381)</td>
<td>62</td>
<td>29</td>
<td>91</td>
<td>3:1</td>
<td>0.13</td>
</tr>
<tr>
<td>F2 (TFRx381)</td>
<td>121</td>
<td>53</td>
<td>174</td>
<td>45:19</td>
<td>0.82</td>
</tr>
<tr>
<td>F2 (SPQx381)</td>
<td>62</td>
<td>29</td>
<td>91</td>
<td>45:19</td>
<td>0.65</td>
</tr>
<tr>
<td>F2 (TFRx381)</td>
<td>123</td>
<td>44</td>
<td>167</td>
<td>3:1</td>
<td>0.70</td>
</tr>
<tr>
<td>F2 (TFRx381)</td>
<td>123</td>
<td>44</td>
<td>167</td>
<td>45:19</td>
<td>0.32</td>
</tr>
<tr>
<td>BC1-F2 (64-F2xSPQ)</td>
<td>28</td>
<td>32</td>
<td>60</td>
<td>1:1*</td>
<td>0.88</td>
</tr>
<tr>
<td>BC1-F2 (47-F2xSPQ)</td>
<td>21</td>
<td>28</td>
<td>49</td>
<td>1:1*</td>
<td>0.61</td>
</tr>
<tr>
<td>BC1-F2 (28-F2xSPQ)</td>
<td>36</td>
<td>29</td>
<td>65</td>
<td>1:1*</td>
<td>0.76</td>
</tr>
<tr>
<td>BC1-F2 (64-F2xSPQ)</td>
<td>28</td>
<td>32</td>
<td>60</td>
<td>1:1</td>
<td>0.88</td>
</tr>
<tr>
<td>BC1-F2 (47-F2xSPQ)</td>
<td>0</td>
<td>49</td>
<td>49</td>
<td>1:1</td>
<td>0.00</td>
</tr>
<tr>
<td>BC1-F2 (28-F2xSPQ)</td>
<td>17</td>
<td>48</td>
<td>65</td>
<td>1:1</td>
<td>0.0004</td>
</tr>
</tbody>
</table>
The statistical analysis of chi-square and percent of co-segregation between the frequency of the single SNP markers in each populations and the phenotype were also performed.

In the two F2 populations, the expected ratios for each SNPs were 3:1 considering the AA and Aa as resistant and aa susceptible. The obtained \( \chi^2 (P) \) of F2 ‘TFRx381e’ SNP1, SNP2 and SNP3 were 0.93, 0.19 and 0.15 respectively. In the F2 ‘SPQx381e’ instead were \( \chi^2 (P) = 0.95 \) for SNP1, \( \chi^2 (P) = 0.59 \) for SNP2 and \( \chi^2 (P) = 1.00 \) for SNP3. In the BC1-F2 was calculated a \( \chi^2 (P) > 0.5 \) in all SNPs analyzed.

In all the populations, the co-segregation between SNP1 marker and phenotypic data was higher of 80% except for the SNP1 valuation in 64-F2, indicating a strong relationship between the SNP1 and the ZYMV resistance genes. In particular, the analysis highlighted the SNP1 could be linked to the major candidate gene and the SNP2 and SNP3, showing a lower percent of co-segregation, to minor genes involved (Table 3.3).

Table 3.3: Based on the associated markers data, chi-square test was calculated for 3:1 expected ratio in the F2 populations and for 1:1, 1:0 and 0:1 expected ratio in BC1-F2 populations. Again, the percent of co-segregation of SNPs vs phenotype was calculated.

<table>
<thead>
<tr>
<th>ID_Population</th>
<th>SNP_ID</th>
<th>Linked markers</th>
<th>No. dominant alleles (A-)</th>
<th>No. recessive alleles (aa)</th>
<th>No. pt total</th>
<th>Expected ratio (R:S)</th>
<th>( \chi^2 (P) )</th>
<th>Percent co-segregation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F2 (TFRx381)</td>
<td>SNP1</td>
<td>Zym1</td>
<td>130</td>
<td>44</td>
<td>174</td>
<td>3:1</td>
<td>0.93</td>
<td>89</td>
</tr>
<tr>
<td></td>
<td>SNP2</td>
<td>Zym2</td>
<td>138</td>
<td>36</td>
<td>174</td>
<td>3:1</td>
<td>0.19</td>
<td>64</td>
</tr>
<tr>
<td></td>
<td>SNP3</td>
<td>Zym3</td>
<td>138</td>
<td>35</td>
<td>173</td>
<td>3:1</td>
<td>0.15</td>
<td>62</td>
</tr>
<tr>
<td>F2 (SPQx381)</td>
<td>SNP1</td>
<td>Zym1</td>
<td>68</td>
<td>23</td>
<td>91</td>
<td>3:1</td>
<td>0.95</td>
<td>80</td>
</tr>
<tr>
<td></td>
<td>SNP2</td>
<td>Zym2</td>
<td>66</td>
<td>25</td>
<td>91</td>
<td>3:1</td>
<td>0.59</td>
<td>72</td>
</tr>
<tr>
<td></td>
<td>SNP3</td>
<td>Zym3</td>
<td>66</td>
<td>22</td>
<td>88</td>
<td>3:1</td>
<td>1</td>
<td>54</td>
</tr>
<tr>
<td>BC1-F2 (64-F2xSPQ)</td>
<td>SNP1</td>
<td>Zym1</td>
<td>28</td>
<td>32</td>
<td>60</td>
<td>1:1</td>
<td>0.61</td>
<td>78</td>
</tr>
<tr>
<td></td>
<td>SNP2</td>
<td>Zym2</td>
<td>68</td>
<td>0</td>
<td>68</td>
<td>1:0</td>
<td>1</td>
<td>28</td>
</tr>
<tr>
<td></td>
<td>SNP3</td>
<td>Zym3</td>
<td>68</td>
<td>0</td>
<td>68</td>
<td>1:0</td>
<td>1</td>
<td>28</td>
</tr>
<tr>
<td>BC1-F2 (47-F2xSPQ)</td>
<td>SNP1</td>
<td>Zym1</td>
<td>21</td>
<td>28</td>
<td>49</td>
<td>1:1</td>
<td>0.32</td>
<td>80</td>
</tr>
<tr>
<td></td>
<td>SNP2</td>
<td>Zym2</td>
<td>0</td>
<td>51</td>
<td>51</td>
<td>0:1</td>
<td>1</td>
<td>57</td>
</tr>
<tr>
<td></td>
<td>SNP3</td>
<td>Zym3</td>
<td>0</td>
<td>51</td>
<td>51</td>
<td>0:1</td>
<td>1</td>
<td>57</td>
</tr>
<tr>
<td>BC1-F2 (28-F2xSPQ)</td>
<td>SNP1</td>
<td>Zym1</td>
<td>36</td>
<td>29</td>
<td>65</td>
<td>1:1</td>
<td>0.39</td>
<td>86</td>
</tr>
<tr>
<td></td>
<td>SNP2</td>
<td>Zym2</td>
<td>31</td>
<td>33</td>
<td>64</td>
<td>1:1</td>
<td>0.80</td>
<td>45</td>
</tr>
<tr>
<td></td>
<td>SNP3</td>
<td>Zym3</td>
<td>29</td>
<td>36</td>
<td>65</td>
<td>1:1</td>
<td>0.39</td>
<td>51</td>
</tr>
</tbody>
</table>
Results obtained from the CAPS analysis also fit with a 3:1 ratio (Table 3.2), confirming a strong relationship between SNP1 and Zym1 resistance genes ($\chi^2 = 0.16, P = 0.7$).

To calculate the overall relationship between phenotypic and genotypic results, we used the Mantel test (Mantel, 1967) on data obtained from ‘TFRx381e’ and ‘SPQx381e’ populations, as shown in figure 3.5. In both populations, the p-value < 0.0001 was lower of significant level alfa = 0.05. The p-value was calculated using the distribution of $r(AB) = 0.186$ in the ‘F2 TFR’ and $r(AB) = 0.196$ in the ‘F2 SPQ’, estimated on 10,000 permutations, allowing to confirm the correlation existing between dissimilarity matrices originated by phenotypic and genotypic dataset of the two F2 segregating populations. Dissimilarity matrices, based on the Euclidean distance, were than used to produce a graphical representation of relationships existing among samples show in figure 3.5.

**Figure 3.5:** Samples clustering, based on phenotypic dataset, in three classes of evaluation of ‘TFRx381e’ (A) and ‘SPQx381e’ (D). Samples clustering, on genetic dataset, in the 23 out of 27 expected allelic configuration segregation of three genes in ‘TFRx381e’ (B) and in ‘SPQx381e’ (E). Mantel test output obtained from the correlation between phenotypic and genotypic matrices.
3.1.5 Analysis of genome scaffolds containing ZYMV markers

The main goal of our work was to develop SNP markers linked to the genes conferring ZYMV resistance. In order to identify genes physically closer to our markers, a functional prediction of protein containing domains involved in the resistance mechanisms was performed. Firstly, the three unigenes containing SNP1, SNP2 and SNP3 identified through the GoldenGate assay have been searched on the genome v2. Table 3.4 reports the scaffolds to which unigenes maps, the percentage of identity and of coverage.

<table>
<thead>
<tr>
<th>Unigene_ID</th>
<th>Scaffold_ID</th>
<th>Identity</th>
<th>Coverage</th>
<th>Start</th>
<th>stop</th>
</tr>
</thead>
<tbody>
<tr>
<td>unigene8819</td>
<td>scaffold1377</td>
<td>97.1%</td>
<td>37.10%</td>
<td>254671</td>
<td>254367</td>
</tr>
<tr>
<td>unigene11643</td>
<td>scaffold1174</td>
<td>99.6%</td>
<td>80.85%</td>
<td>546127</td>
<td>546865</td>
</tr>
<tr>
<td>unigene11643</td>
<td>scaffold1174</td>
<td>98.4%</td>
<td>19.80%</td>
<td>544665</td>
<td>544847</td>
</tr>
<tr>
<td>unigene9577</td>
<td>scaffold2494</td>
<td>99.5%</td>
<td>100.00%</td>
<td>79461</td>
<td>80079</td>
</tr>
</tbody>
</table>

The scaffolds with the highest percentage of identity and coverage is the scaffold2494 for the unigene containing the SNP1, scaffold1377 for the unigene of SNP2 and the scaffold1174 for SNP3. The length of these three scaffolds was between 257.375 bp and 714.477 bp. GenScan tool was used to translate the scaffolds sequences in the peptides in order to interrogate by InterProScan tool. The InterProScan prediction carried out on scaffolds allow us to find a closely association of the SNP1 with two CNL domains, a RLK domain and an ATP-dependent RNA helicase as shown in figure 3.6. ATP-dependent RNA helicase, LRR and other domains potentially involved in pathogen recognition were also found on the scaffolds to which belong the SNP2 and SNP3.
**Figure 3.6:** By InterProScan analysis, SNP1 marker results to be associated to ATP-dependent RNA helicase, two CNL and a RLK (data not showed) in genome v2.

The close relationship found between the R-domains and SNP1 suggest us that such marker could be associated to one of three resistance genes.
3.2 Transcriptome analysis of zucchini reference genome and of susceptible and resistant lines

3.2.1 Reference transcriptome assembling and functional annotation

In order to perform a deep functional annotation of *C. pepo* reference transcriptome, the version 3 was assembled in 108,062 transcripts, revealing 90% of overall alignment of transcripts over the genome. Out of 108,062 reference transcripts 79,733 showed a full transcribed sequence and 28,330 fail to give a complete transcript. They were classified as not full transcripts and further explored in order to identify broken sequences at scaffold ends. Approximately 1000 cases were extracted and added to first dataset because full transcript was reconstructed.

A total of 108,062 transcripts were analyzed using the Blast2GO tool (B2GO; Conesa et al., 2005), at the default parameters, in order to understand their putative functions. The analysis revealed that 4.44% of the transcripts do not showed similarity with known sequences collected in interrogated databases. On the other hand, only the 36.54% of the remaining transcripts (95%) were classified according the terms included in the three main ontology categories (molecular function, biological process and cellular component) proposed by the Gene Ontology consortium which provides an ontology of defined terms representing gene product properties. A relevant portion transcripts (39,486) showed a partial human readably annotation. The data distribution of B2GO analysis is showed in figure 3.11.

![Figure 3.11: Distribution of B2GO analyzed transcripts belongs to reference transcriptome v3.](image-url)
This analysis was performed in order to identify the biological function of each zucchini transcript. In figure 3.12 is reported the graphical representation of the distribution of GO terms among the three ontologic category (P) biological process, (F) molecular function and (C) cellular component, based on 108,062 transcripts. In the biological process, the two biggest functional categories were represented by the “metabolic process” and “cellular process”, with 20,666 and 19,588 transcripts respectively. Out of annotated transcripts in the molecular function category, 19,648 transcripts were involved in “binding” and 17,512 transcripts in “catalytic activity”. In the cellular component, the three most representative functional categories were “cell” (13,671), “organelle” (9,432) and “membrane” (7,175).

![Figure 3.12: Transcripts level 2 classification in (A) Biological process, (B) molecular function and (C) Cellular component ontology categories.](image-url)
3.2.2 Transcriptome sequencing and mapping of zucchini resistant and susceptible genotypes

RNA sequencing experiment was performed on three biological replicates of two resistant and susceptible genotypes inoculated with ZYMV collected at 12 and 22 dpi. A paired-end tag sequencing strategy was chosen, in which short and paired tags were extracted from the ends of long DNA fragments for ultra-high-throughput sequencing. The protocol included strand-specific library preparation followed by 100 base pair (bp) paired-end reads sequencing. Before to perform further analysis, a quality check was conducted on the raw sequencing data, removing low quality portions and preserving the longest high quality part of a NGS read. RNA-seq generated around 36 million reads of 100 bp length.

In order to ensure a high quality and reliability of the analysis was established a minimum sequence length of 25 bp and a sequence quality score of 35. The total number of reads per sample before and after the trimming, the number of sequenced bases and the coverage mapping is reported in table 3.6.

<table>
<thead>
<tr>
<th>Library type</th>
<th>«381e»</th>
<th>«TFR»</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paired ends</td>
<td>Paired ends</td>
<td></td>
</tr>
<tr>
<td>Sequenced reads length</td>
<td>2 x 100 bp</td>
<td>2 x 100 bp</td>
</tr>
<tr>
<td>Number of reads before trimming</td>
<td>163,483,878.00</td>
<td>126,621,462.00</td>
</tr>
<tr>
<td>Number of reads after trimming</td>
<td>151,087,631.00</td>
<td>112,382,770.00</td>
</tr>
<tr>
<td>Sequenced bases</td>
<td>30.22 Gbp</td>
<td>22.48 Gbp</td>
</tr>
</tbody>
</table>

The high quality reads, obtained after trimming, were aligned against the Cucurbita pepo transcriptome v3, assigning a number of reads to each annotated locus and calculating the relative abundances. The resulting alignment files were used as input for the calculation gene expression values. The output file includes one column for the transcript ID and one column for expected-read-counts for each sample. Columns correspond to the three biological replicates for the four conditions under analysis (‘True French’ and ‘381e’ collected both at 12 and 22 dpi). Data counts were normalized using the RPKM method (Mortazavi et al. 2008).

The quality control of the mapped reads of ‘381e’ and ‘TFR’ genotypes using TopHat, allow us to remove the mapped reads with quality score < 30 and remove duplicates generated during the sequencing reaction. The two Quality BAM files produced were graphically represented in figure 3.8.
Figure 3.8: Graphical representation of number of alignments in various mapping quality (MAPQ) intervals and number of mapped sequences of ‘381e’ genotype (A) and ‘TFR’ genotype (B).

The assembled transcripts, 73348 and 73609 for ‘TFR’ and ‘381e’ respectively, have a mean length of 1052 bp, minimum length of 201 bp and a maximum length of 9917 bp. The overall quality of the biological replicates was evaluated, based on the similarity between replicates, by a PCA analysis. At first, the similarity was calculated using obtained reads reported in table 3.6. Since the zucchini available transcriptome version 3 is very fragmented and the annotation does not distinguish between splicing variants and transcripts of different loci. For this reason, were extracted the unique transcripts with expression values different from the others and for which the reads mapped univocally. Therefore, the 32,520 unique transcripts were selected. The quality control and the samples clustering is shown in figure 3.9.
Figure 3.9: Clustering of sequenced samples using all the reads (A) and only the unique transcripts obtained after the repeated quality control (B).

The similarity between replicates and the Euclidean distance between the samples was evaluated by hierarchical clustering. Using this approach, we ascertain if any samples behave differently by the other replicates (outliers). The cluster analyses highlighted that some replicates did not behave as expected showing too much divergence from the other replicates of the same group. Therefore, samples 38IA12, 38IA22 and TFC12 were removed from the downstream analysis. As the TF-22 replicates were the most uniform, all of them were used for the following analyses. The resulting plots, performed with the 32,520 unique transcripts, before (A) and after (B) the removal of the outliers are showed below in the figure 3.10.

Figure 3.10: Unique transcripts (32,520) clustering obtained after the repeated quality control of all sequenced samples (A) and after removed outliers (B).
Because of the low quality of the available reference transcriptome, the following step was to annotate the transcriptome to the genome, in order to obtain the coordinates of the transcripts/genes over the genome, before starting the RNA-seq analysis.

### 3.2.3 Identification of Differential Expressed Genes

Differential expression analysis has been performed on the final datasets without outliers. To discover the differentially expressed genes (DEGs), the following comparisons were included in the analysis:

- 381-12dpi vs TF-12dpi
- 381-22dpi vs 381-12dpi
- 381-22dpi vs TF-22dpi
- TF-22dpi vs TF-12dpi
- 381 (all times) vs TF (all times)

A change was deemed significant based on the following criteria: the transcript is scored and the q-value of the statistical test is less than 0.05. In order to get the list of the mostly interesting genes, a further filter was applied by selecting those with a fold change higher than 1.5-fold (UP-regulated) and lower than -1.5-fold (DOWN-regulated). The 381-12dpi vs TF-12dpi comparison aims to the identification of the transcripts differentially expressed (DE) in the ‘381e’ resistant line respect to ‘True French’ susceptible line after 12 day from virus inoculation. A total of 680 up-regulated and 900 down-regulated genes were identified in the resistant line.

The 381-22dpi vs 381-12dpi comparison aims the identification of the transcripts differentially expressed (DE) in the 381-22dpi line respect to 381-12dpi line. Out of a total 1302 total DE transcripts, 656 were up-regulated and 646 down-regulated.

The 381-22dpi vs TF-22dpi comparison aims the identification of the transcripts differentially expressed (DE) in the 381-22dpi line respect to TF-22dpi line. A total of 484 transcripts were found up-regulated in 381-22dpi line respect to TF-22dpi line and 533 down-regulated.

The TF-22dpi vs TF-12dpi comparison aims the identification of the transcripts differentially expressed (DE) in the TF-22dpi line respect to TF-12dpi line. A total of 1164 transcripts were up-regulated in TF-22dpi line respect to TF-12dpi line and 1088 down-regulated.
In addition to the comparisons already described, a more general analysis has been performed in order to identify DE transcripts in ‘381e’ genotype respect to ‘TFR’ genotype. In this experiment, the time was not treated as factor. This comparison aims the identification of the transcripts differentially expressed (DE) in the ‘381e’ genotype samples respect to ‘TFR’ genotype samples. In this comparison, a total of 350 transcripts were up-regulated in ‘381e’ genotype respect to ‘TFR’ genotype and 641 down-regulated.

In order to identify the transcripts that are significantly expressed across the selected experimental conditions, the expressed genes for both genotype in both sampling times were recorded. Comparison of common and specific genes express in each studied condition was performed (Figure 3.13 A). A total number of 1,302 expressed genes were found for ‘381e’ resistant genotype, while ‘TFR’ susceptible genotype showed 2,252 expressed genes, regardless the sampling time. ‘381e’ presented 1,006 expressed genes absent in ‘TFR’ as well as ‘TFR’ presented 1,956 expressed genes absent in ‘381e’, while there were 296 common genes (Figure 3.13 B).

It is interesting to note, that in both diagram there is a high number of specific expressed genes and, in contrast, there is a reduced number of common genes for each sample, which may be related to the sampling time of inoculum.

**Figure 3.13:** Venn diagram showing the number of overlapping and non-overlapping expressed genes in the comparisons between the two sampling time of inoculum 12 and 22 dpi (A) and two genotypes (B).
Studying more thoroughly the unique genes expressed in the two genotypes (1006 and 1956), several pathogenesis-related proteins were identified. Those showed a relative abundance of transcripts coding for LRR domain and Serine/threonine kinase in both genotypes, meanwhile 9 ATP-dependent helicases was detected in the resistant genotype with respect the 6 found in susceptible genotype. More details were showed in figure 3.14.

![Pathogenes-related proteins and domains](image)

**Figure 3.14**: Graphical representation of pathogenesis-related protein and domains identified in 1006 and 1956 genes expressed uniquely in ‘381e’ and ‘TFR’ genotypes respectively.

### 3.2.4 R-genes prediction

The *Cucurbita pepo* proteome was also interrogated by Matrix-R tools that, via HMM profiling, in order to retrieve, annotate and classify plant resistance genes. The protein sequences of the genes identified, were examined in order to underscore typical R gene domains. The output was a list of transcripts with a hit to a resistance domain. Out of 45016 only 2113 transcripts (belongs to the *C. pepo* reference transcriptome) have a score of "HIT" for at least one R-domain.
3.2.5 SNP calling analysis

In order to discover new SNP markers associated to genes involved in the defense mechanisms, a SNP calling analysis against the *Cucurbita pepo* reference genome was performed for transcripts of the susceptible ‘TFR’ and resistant ‘381e’ accessions. This analysis, allows to highlight the total number of 30275 and 6150 SNPs for ‘381e’ and ‘TFR’ respectively (figure 3.15). The lists of genes carrying SNPs obtained by SNP calling analysis were compared with the lists of genes up and down regulated of two genotypes highlighting the number of DE genes having SNPs as showed in Venn diagram reported in figure 3.11. In particular out of 350 up-regulated genes in ‘381e’, 56 contained SNPs, as well as of 641 down-regulated genes of ‘381e’ genotype, 121 carried least a SNP in comparison with the reference *C. pepo* genome. Likewise, out of 641 up-regulated and 350 down-regulated genes in ‘TFR’ genotype 48 and 22 respectively, contain at least a SNP.

![Venn diagram showing the number of ‘TFR’ and ‘381e’ DEGs having SNPs compared to the reference *C. pepo* genome.](image)

Out of total DE genes in ‘TFR’ and ‘381e’ carrying least a SNP marker, a high percentage coding for protein kinase (42 and 27), Chloroplast protein kinase (28 and 26), Leucine Reach Repeats (10 and 1) and other important protein such as Thaumatin, Ubiquitin, Phosphatase, but also hormones such as Auxin and Ethilene, typically involved in response reaction of plants to biotic stresses was found (Figure 3.16).
Figure 3.16: Graphical representation of pathogenesis-related protein identified in up and down regulated genes of ‘381e’ and ‘TFR’ genotypes carrying least a SNP marker.

Inconsistent and low quality/coverage SNPs were removed, identifying 3357 unique SNPs for ‘TFR’ and 14327 unique SNPs for ‘381e’ enlarged the number of SNPs markers available for C. pepo (Figure 3.17) and enhancing our possibility to find new markers associated with resistance to ZYMV. Interestingly, the presence of SNPs putatively associated to genes Zym-1, Zym-2 and Zym-3, previously analyzed by the HRM assay on five different segregating populations was confirmed, demonstrating that they can be good candidate markers for resistance. Particularly, the SNP1 and SNP2 belongs to the 14327 SNPs of ‘381e’ genotype, while the SNP3 ranged in the 3357 SNPs of ‘TFR’ genotype.

Next step was to select a pool of SNPs, targeting genes, between the two susceptible and resistant isogenic lines (Figure 3.17). The SNP calling was performed on the transcriptome of the two genotypes taking into account only the expressed genes in a given time, excluding from the assessment all the SNPs belongs the not expressed genes and the intronic SNPs. This procedure excluded intron alleles that can affect disease such as transcription regulators, leaving analysis to genome elements of large effect.
3.2.6 Identification of new candidate markers for ZYMV resistance

In order to identify new candidate markers for ZYMV resistance we merged SNP calling and Matrix-R data. Unique SNPs for each genotype were filtered in order to extract only SNPs located on resistance-related transcripts highlighted by Matrix-R analysis. Results showed that out of 3357 and 14327 unique SNPs identified in ‘TFR’ and ‘381e’ genotypes respectively, 251 SNPs and 1895 SNPs fall in R genes or in domains involved in defense mechanisms. Moreover, once a SNP was placed on a scaffold, we were able to identify putative pathogen recognition genes localized in the same chromosome region, allowing us to focus on a restricted set of genes. A functional R gene map based on the co-localization of pathogen recognition putative proteins with R loci linked markers on scaffold 2494 is shown in figure 3.18. The map shows the candidate pathogen recognition genes localized on scaffold and the distribution of ‘381e’ genotype SNPs.
The visualization of information linked to a locus is a fundamental step in interpreting data and in suggesting correlations between genetic and genomic data. Indeed, 100 SNPs of ‘381e’ are falling in only three R genes located on scaffold 2494 (figure 3.19 A), the same scaffold to which belongs the SNP1 associated to major gene Zym-1. In parallel, out of 251 SNPs discovered in the ‘TFR’ genotype, 19 falling in three R genes of scaffold 1174 (figure 3.19 B), that holds also SNP3 putatively associated to Zym-3. While, we not found SNPs falling in R genes located on scaffold 1377 where SNP2 was identified. In addition, it is important to note that the distribution of SNP markers allowed us to identify a peak of SNPs in both genotypes in scaffold 1518 with a higher number in the genotype ‘381e’ (159 SNPs) versus genotype ‘TFR’ (34 SNPs).
Figure 3.19: R genes-related SNPs distribution on genome scaffolds (SNPs vs scaffolds genome). On the scaffold2494 (green bar in panel A) 100 SNPs of ‘381e’ falling on only three R genes. On the scaffold1174 (red bar in panel B) 19 SNPs of ‘TFR’ falling on three R genes.
4. DISCUSSIONS

The use of resistant varieties in agriculture provides both economic and ecological benefits, considering the reduced use of pesticides needed for their cultivation. The process of introduction of resistance to pathogens in crop species, particularly those polygenic, takes long time and are not always stable over time. Molecular genetics and modern biotechnology techniques can be very useful to identify pathogen resistance genes in related wild species and to speed up the introgression in cultivated varieties. Several studies rely on use of advanced segregating populations for accelerating the introgression of genes of interest into a genotype. This process is also called gene pyramiding, defined as a method aimed to assembling multiple desirable genes from multiple parents into a single genotype (Ye and Smith, 2008).

Main goal of this work was to study the association of molecular markers with resistance genes to ZYMV, in order to make easier, faster and economically sustainable the introgression of these genes in traditional Campania variety ‘San Pasquale’. The identification of genetic resource containing the pathogen resistance genes and the knowledge of mode of inheritance is a very important task for carrying out a breeding program. For this purpose, several in vivo assays were conducted to evaluate parental lines behavior and to estimate segregation ratio of genes involved in resistance to ZYMV in progenies.

The phenotypic in vivo assays, performed on two populations ‘F₂ TRF’ and ‘F₂ SPQ’ and three derived ‘F₂ SPQ’ backcross populations, allowed us to screen resistant and susceptible samples to a ZYMV breaking strain found in the major growing regions of the crop. In vivo tests was conducted in environmental controlled conditions in order to ensure a high stress condition. The F₂ segregation populations displayed the 45:19 expected ratio, obtained assuming three complementary dominant genes involved in resistance to ZYMV, reported in previous segregating populations (Paris and Cohen, 2000). However, the phenotypic assays performed on BC₁-F₂ populations not respected the expected ratio based on Paris and Cohen (2000) assumptions. It is important to note that the genotypes carrying AAbbcc and Aabbbc alleles results resistant. If we hypothesized a major gene control the resistance the observed ratio fitted with the 1:1 expected ratio in the BC₁-F₂ populations. Following this hypothesis we recalculated the segregation in the F₂ populations showing to be in agreement also with 3:1 ratio. This finding highlighted a very important role of Zym-1 in conferring resistance. The two complementary Zym-2 and Zym-3 genes could be nevertheless indispensable for ensuring the expression of tolerance in late stages of development. In the study of resistance to TMV and Phytophthora capsici in pepper, important epistatic effects between a major and three minor QTLs (with or without additive effects) were detected (Lefebvre and Palloix, 1996).
The main role of Zym-1 gene was also confirmed by the fact that the percent of co-segregation with the phenotype was lower when the genotypes having only the Zym-1 was considered susceptible especially in the BC1-F2 populations (Table 3.2) and by Mantel test (Mantel, 1967) conducted on the (dis)similarity or distance matrices obtained from two F2 segregating populations datasets. Molecular analysis performed with the High Resolution Melting Real-Time PCR, based on each single SNP marker linked to the three resistance genes, allowed us to predict the allelic configurations of the considered loci. Moreover, statistical analysis based on the SNPs dataset showed a match between the observed and expected ratio of segregation in four segregating populations, confirming a higher co-segregation of Zym-1 with the resistant phenotype respect to minor genes. The CAPS marker analysis discriminating susceptible from resistant subjects, confirmed the results obtained from the HRM analysis showing a stronger co-segregation of SNP1 with resistant phenotypes.

The SNPs and CAPS markers obtained in the study enriched molecular markers of C. pepo. CAPS markers are widely used in molecular assisted selection programs and in diagnostic assays of vegetable crops. The relevance of a CAPS marker for MAS was demonstrated in different breeding programs (Langella et al., 2004; Suh et al., 2013; Joshi et al., 2013). The rapidity, accuracy and the cost efficiency of our CAPS analysis, makes it a useful tool to improve the traditional selection methods of breeding.

Molecular markers develop in this thesis can help future genetic linkage and quantitative trait loci analysis, and will be essential to speed up the process of breeding new and better adapted squash varieties (Blanca et al., 2011). This work highlighted three SNP markers linked to three genes putatively responsible of the ZYMV resistance. The ZYMV resistance genetic segregation studied both at phenotypic and genotypic level not showed a clear framework of action of genes involved. Therefore, a different mode of inheritance or the presence of additional genes involved in resistance process could be required. For example, Pachner et al. (2014) theorizes a combined dosage effect of the genes that conferring determinate phenotypes. The dosage effect of genes could explain the recovery cases after 25th dpi of same analyzed samples. Gomez et al. (2009) stated that 51% of resistance traits to plant viruses were dominant, 35% were recessive, and the remainder were more complex (incomplete dominance or dose-dependent).

Marker-based selection reduces/eliminates extensive phenotyping, reducing the selection times, excessive costs due to the large spaces needed for plants and pathogens growth, but also eliminating the errors owed to visual assessments often wrong because made subjectively by the operator. The rate of success is likely to increase due to developments in gene-based marker development, more
efficient quantitative trait locus mapping procedures, and lower cost genotyping systems (Xu and Crouch, 2008). However, in many cases, particularly regarding complex traits, it is difficult to assign a genotype to specific symptoms. Therefore, the most reasonable approach towards merging the various genes for resistance into one genotype is a step-wise combination of phenotypic and marker-assisted selection (Pachner et al., 2014). Development of high throughput precision phenotyping systems for resistance gene mapping, improved understanding of genotype by environment interaction and epistasis, and development of publicly available computational tools tailored to the needs of molecular breeding programs are required (Xu and Crouch, 2008).

Genomic analysis appears to be increasingly useful in the associated markers discovery and resistance genes study. In order to deeply investigate genes-markers association several modern bioinformatics tools were used. To obtain a better overview of R genes co-localized next to our markers we performed a protein prediction and domain search through InterProScan tool (Andolfo et al., 2014). Our attention was focused to the research of protein belonging to resistance gene classes. Proteins belonging to CC, NB-ARC, LRR, ATP-dependent RNA helicase and other typical R-domains were found nearby the SNP1 markers, suggesting the SNP1 association with one of three resistance genes, most probably the major gene Zym-1. R genes and/or gene silencing mediated active resistance mechanisms. The NB-ARC system minimizes the cost of defense for the plant, as multiple NB-ARC-LRR proteins can be maintained at a low level in the absence of a pathogen and rapidly induced under pathogen attack through miRNA regulation (Li et al., 2011). Pathogen-encoded suppressors of RNA-silencing mechanisms might result in the induction of multiple NB-ARC-LRR defense proteins (Shivaprasad et al., 2012). The RNA helicase are enzymes that use energy derived from the hydrolysis of a nucleotide triphosphate to unwind double-stranded RNAs (de la Cruz et al., 1999). RNA helicases have been shown to be involved in every step of RNA metabolism, including nuclear transcription, pre-mRNA splicing, ribosome biogenesis, nucleocytoplasmic transport, translation, RNA decay, and organellar gene expression (Tanner et al., 2001). Xu et al. (2013) reported the involvement of RNA helicases in response to stress and in plant growth and development.

Plant-expressed molecules during pathogen challenging can give insights into the underlying defense mechanisms. Plants have evolved a defense system against pathogens that involves the regulation of gene expression, cascade signaling activation, hormone balancing and synthesis of defensive metabolites (Mithöfer and Boland, 2012). The advent of next-generation sequencing (NGS) has revolutionized genomic and transcriptomic approaches to biology. These new
sequencing tools are also valuable for the discovery, validation and assessment of genetic markers in populations (Davey et al., 2011).

In order to discovery new genes and their associated markers involved in defense mechanisms to ZYMV in *Cucurbita pepo*, the transcriptomes of one resistant and one susceptible genotype collected in two different time post-inoculation with ZYMV were sequenced. Through the RNA-seq analysis, transcriptional changes were monitored in zucchini upon inoculation with virus. Several researches in this field were conducted in order to identify genes connected to resistance (Li et al., 2013; Sookruksawong et al., 2013; Xue et al., 2013; Zeng et al., 2013). RNA-seq technology, it has also been proving to be an extremely useful tool for identifying single nucleotide polymorphisms, which may be also used to develop high density SNPs chips for studies concerning the genome wide association (GWAS) and to build high density linkage maps (Baranski et al., 2014; Yu et al., 2014). Furthermore, SNPs can be used as markers in order to distinguish allelic transcripts whilst studying the allele-specific expression (Bell and Beck, 2009).

At our knowledge, this is the first work actually available reporting study of zucchini genotypes response to ZYMV attacks through RNA-seq assay. The major limit to analyze transcriptomic data is due to the very fragmented and poorly annotated reference transcriptome of *C. pepo* currently available. Therefore, in this work has been carried out the functional annotation of reference transcriptome version 3, allowing us to obtain more detailed information useful for B2GO annotation of differentially induced genes in our analysis. In genomes that have been poorly studied, sequence availability is the limiting factor for the discovery of SNPs (Blanca et al., 2011). The functional annotation of unique genes expressed in the two genotypes (1006 and 1956), allowed us to identify several transcripts coding for LRR domains and Serine/threonine protein kinase in both genotypes, meanwhile a higher presence of ATP-dependent helicases was detected in the resistant genotype, enlarging the number of SNPs available for this species.

In order to discovery new markers and R genes, a SNP calling and Matrix-R analysis were performed on the sequenced transcriptome of resistance and susceptible genotypes, detecting the genotype-specific SNPs and R genes related SNPs. Merely in the 2011, Blanca *et al.* conducted the first work of SNPs discovery using Expressed sequenced tags (ESTs). SNP calling in the context of NGS data analysis is performed against reference genome, typically including an associated confidence score or statistical evidence metric (Li *et al.*, 2008).

In our work SNP calling was performed on the transcriptome of the two genotypes taking into account only the expressed genes in a given time, excluding from the assessment all the SNPs belongs the not expressed genes and the intronic SNPs. This procedure excluded intron alleles that can affect disease such as transcription regulators, leaving analysis to genome elements of large
effect. While some correlation exists between exon to intron variation, only whole genome sequencing would be able to capture the source of all relevant SNPs. The SNP calling allowed us to detect unique SNPs comparing the two transcriptomes against the reference genome of *Cucurbita pepo*. Stringent quality criteria were used for distinguish sequence variations from sequencing errors and mutations introduced during the cDNA synthesis step. By applying these criteria were identified unique SNPs both for ‘TFR’ and ‘381e’, enhancing our possibility to find new resistance genes associated markers.

R genes prediction with Matrix-R assay, based on the reference transcriptome B2GO annotation, allowed to predict genes putatively involved in defense response. Merging information obtained from SNP calling and Matrix-R assay, we extracted the unique SNPs related to R-gene, discovering a high SNPs concentration (peak) in correspondence of genome portion where are located Zym-1 and Zym-3 genes. Detected SNPs could be defined “putatively ZYMV resistance associated”, considering that the analysis was performed on resistant and susceptible isogenic lines dataset. Interestingly a further peak of SNPs in both genotypes was identified in scaffold1518 although in higher numbers in the genotype ‘381e’.

Gene expression differences highlighted in this work between the resistant and susceptible isogenic lines was used for a comprehensive study of expressed genes and for discover expressed SNPs in zucchini in response to ZYMV. Comparing the lists of genes containing SNP markers with lists of genes of ‘TFR’ and ‘381e’ induced or repressed by the virus attack, were highlighted genes differentially expressed in the two genotypes having one or more polymorphisms. The identification and functional annotation of polymorphic genes identified represent a useful resource to comprehend mechanisms determining complex traits and, consequently, to develop programs aiming the genetic improvement (Baranski *et al.*; Yu *et al.*, 2014).
5. CONCLUSIONS

Resistant varieties produce significant economic and ecological benefits by reducing inputs of pesticides and, therefore, their use is recommend in integrated and biological cultivation systems. The process of disease resistance transfer by classical breeding in the cultivated varieties requires long times, some resistance genes are difficult to introduce and are not always stable over time. The use of molecular markers associated to the genes controlling resistance characters may facilitate the selection.

In this work a study of ZYMV resistance response in zucchini at genetic and genomic level was performed.

- Phenotypic and genotypic analysis, performed on five segregating populations, highlighted a complex inheritance of resistance investigated with the involvement of at least three genes (Zym-1, Zym-2 and Zym-3) in the defense mechanisms. The analyses showed that Zym-1 has a main role in the zucchini response to virus and that Zym-2 and Zym-3 could be required for displaying a durable resistance in later stages of plants development.

- Molecular segregation analysis on 3 SNP loci strongly suggested a relationship between these SNPs and resistance genes Zym-1, Zym-2 and Zym-3. A CAPS marker developed on SNP1 validated the association between the SNP1 and the Zym-1 resistance gene and it is available to improve selection methods employed at breeding companies.

- Transcriptomic analysis allowed to observe above 3000 differentially expressed genes between zucchini isogenic lines for resistance to ZYMV highlighting additional resistance gene candidates. Thanks to the functional annotation performed on the DEGs and comparison of the differentially expressed genes an explanatory model of the response process of zucchini to ZYMV could be build up.

- Transcript SNP calling enlarged the number of marker available for C. pepo and reconfirmed the presence of SNPs, putatively associated to genes Zym-1, Zym-2 and Zym-3, in the two isogenic lines ‘True French’ and ‘381e’. SNPs localized near R genes evidenced SNPs associated to typical resistance domain expressed in differential mode in resistant and susceptible genotypes.
• The R protein domains prediction showed a close relationship of the SNP1 with a CNL and an ATP-dependent RNA helicase, suggesting the SNP1 association with one of three resistance genes, most probably the major gene Zym-1.

• Considering that the local variety ‘San Pasquale’, susceptible to ZYMV, has a good marketability in Campania region, results obtained in this thesis can be useful for the genetic improvement of this variety.
6. REFERENCES


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8. STAGE

The stage was conducted from 17 November 2014 to 18 December 2014 at the “SEQUENTIA BIOTECH” laboratory, Campus UAB - Edifici CRAG - Bellaterra - Cerdanyola del Vallès – BARCELONA. During this stage, it was performed research about RNA-seq analysis and functional gene annotation with particular attention to the SNPs discovery and R-genes analysis.

9. PUBLICATIONS


Capuozzo C., Manzo D., Formisano G., Rrcolano M.R. CAPS and HRM markers for rapid detection of ZYMV resistance loci in zucchini. 57° Congresso annuale SIGA – Foggia 16-19 September 2013
