Characterization of the USDA *Cucurbita pepo*, *C. moschata*, and *C. maxima Germplasm* Collections

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The *Cucurbita* genus is home to a number of economically and culturally important species. We present the analysis of genotype data generated through genotyping-by-sequencing of the USDA germplasm collections of *Cucurbita pepo*, *C. moschata*, and *C. maxima*. These collections include a mixture of wild, landrace, and cultivated specimens from all over the world. Roughly 1,500 - 32,000 high-quality single nucleotide polymorphisms (SNPs) were called in each of the collections, which ranged in size from 314 to 829 accessions. Genomic analyses were conducted to characterize the diversity in each of the species and revealed extensive structure corresponding to a combination of geographical origin and morphotype/market class. Genome-wide associate studies (GWAS) were conducted for each data set using both historical and contemporary data, and signals were detected for several traits. These data represent the largest collection of sequenced *Cucurbita* and can be used to direct the maintenance of genetic diversity, for developing breeding resources, and to help prioritize whole-genome re-sequencing.

# Introduction

The *Cucurbitaceae* (Cucurbit) family is home to a number of vining species mostly cultivated for their fruits. This diverse and economically important family includes cucumber (*Cucumis sativus*), melon (*Cucumis melo*), watermelon (*Citrullus lanatus*), and squash (*Cucurbita ssp.*). Like other cucurbits, squash exhibit diversity in growth habit, fruit morphology, metabolite content and disease resistance, and have a nuanced domestication story . The genomes of *Cucurbita ssp.* are small (roughly 400 Mb), but result from complex interactions between ancient genomes brought together through an allopolyploidization event . These factors make squash an excellent model for understanding the biology of genomes, fruit development, and domestication. Within *Cucurbita*, five species are recognized as domesticated. Three of these are broadly cultivated: *C. maxima*, *C. moschata*, and *C. pepo* . Few genomic resources have been available for these species; although, draft genomes and annotations, along with web-based tools and other genomics data are emerging . Already, these resources have been used to elucidate the genetics of fruit quality, growth habit, disease resistance, and to increase the efficiency of cucurbit improvement ; however, there has yet to be a comprehensive survey of the genetic diversity in the large diverse *Cucurbita* germplasm panels maintained by the USDA within the National Plant Germplasm System.

Germplasm collections play a vital role in maintaining and preserving genetic variation. These collections can be mined by breeders for valuable alleles. They can also be used by geneticists and biologists for mapping studies . Like many other orphan and specialty crops,there has been little effort put into developing community genetic resources for squash and other cucurbits. The Cucurbit Coordinated Agricultural Project (CucCAP project) was established to help close the knowledge gap in cucurbits . This collaborative project aims to provide genomics resources and tools that can aid in both applied breeding and basic research. The genetic and phenotypic diversity present in the USDA watermelon, melon, and cucumber collections has already been explored as part of the CucCap project, partially through the sequencing of USDA germplasm collections and development of core collections for whole-genome sequencing . The diverse specimens of the USDA squash collections have yet to be well characterized at the genetic level;although, an elaborate system has been established for classifying squash based on species and various other characteristics.

The classification system used in squash is complex. Squash from each species can be classed as winter or summer squash depending on whether the fruit is consumed at an immature or mature stage, the latter is a winter squash . Squash are considered ornamental if they are used for decoration, and some irregularly shaped, inedible ornamental squash are called gourds; however, gourds include members of *Cucurbita* as well as some species from *Lagenaria*—not all gourds are squash . Many squash are known as pumpkins; the pumpkin designation is a culture dependent colloquialism that can refer to jack O’ lantern types, squash used for desserts or, in some Latin American countries, to eating squash from *C. moschata* known locally as Calabaza . Cultivars deemed as pumpkins can be found in all widely cultivated squash species. Unlike the previous groupings, morphotypes/market classes are defined within species. For example, a Zucchini is reliably a member of *C. pepo* and a Buttercups are from *C. maxima*. Adding to the complexity of their classification, the *Cucurbita* species are believed to have arisen from independent domestication events and the relationships between cultivated and wild species remains poorly understood .

*C. pepo* is the most economically important of the *Cucurbita* species and is split into two different subspecies: *C. pepo* subsp. *pepo* and *C. pepo* subsp. *ovifera* . Evidence points to Mexico as the center of origin for *pepo* and southwest/central United States as the origin of *ovifera*. The progenitor of *ovifera* is considered by some to be subsp. *ovifera* var. *texana*, whereas subsp. *fraterna* is a candidate progenitor for *pepo* . Europe played a crucial role as a secondary center of diversification for subsp. *pepo*, but not subsp. *ovifera* . Important morphoptypes of *pepo* include Zucchini, Spaghetti squash, Cocozelle, Vegetable Marrow, and some ornamental pumpkins. *C. pepo* subsp. *ovifera* includes summer squash from the Crookneck, Scallop, and Straightneck group, and winter squash such as Delicata and Acorn .

The origin of *C. moschata* is more uncertain than *C. pepo*; it is unclear whether *C. moschata* has a South or North American origin . Where and when domestication occurred for this species is also unknown; however, it is known that *C. moschata* had an India-Myanmar secondary center of origin where the species was further diversified . *C. moschata* plays an important role in squash breeding as it is cross-fertile to various degrees with *C. pepo* and *C. maxima*, and can thus be used as a bridge to move genes across species . Popular market classes of *C. moschata* include cheese types like dickenson, which is widely used for canned pumpkin products, Butternut (Neck) types, Japonica, and tropical pumpkins known as Calabaza .

*C. maxima* contains many popular winter squash including Buttercup/Kabocha types, Kuri, Hubbard, and Banana squash . This species also sports the world’s largest fruit, the giant pumpkin, whose fruit are grown for competition and can reach well over 1000 Kg . Although this species exhibits a wide range of phenotypic diversity in terms of fruit characteristics, it appears to be the least genetically diverse of the three species described . *C. maxima* is believed to have a South American origin, and was likely domesticated near Peru, with a secondary center of domestication in Japan and China .

In this study, we set out to characterize the genetic diversity present in the USDA *Cucurbita* germplasm collections for *C. pepo*, *C. moschata*, and *C. maxima*. We present genotyping-by-sequencing (GBS) data from each of these collections, population genomics analysis, results from genome-wide association stud (GWAS) using historical and contemporary phenotypes, and suggest a core panel for re-sequencing.

# Materials and Methods

## Plant Materials and Genotyping

All available germplasm were requested from USDA cooperators for *C. maxima* (534 accessions from Geneva, NY), *C. moschata* (314 accessions from Griffin, GA), and *C. pepo* (829 accessions from Ames, IA) respectively. Seeds were planted in 50-cell trays and two 3/4 inch punches of tissue (approximately 80-150 mg) was sampled from the first true leaf of each seedling. DNA was extracted using Omega Mag-Bind Plant DNA DS kits (M1130, Omega Bio-Tek, Norcross, GA) and quantified using Quant-iT PicoGreen dsDNA Kit (Invitrogen, Carlsbad, CA). Purified DNA was shipped to Cornell’s Genomic Diversity Facility for GBS library preparation using protocols optimized for each species. Libraries were sequenced at either 96, 192, or 384-plex on the HiSeq 2500 (Illumina Inc., USA) with single-end mode and a read length of 101 bp.

## Variant Calling and Filtering

SNP calling was conducted using the TASSEL-GBS V5 pipeline . Tags produced by this pipeline were aligned using the default settings of the BWA aligner . Raw variants were filtered using BCFtools . Settings for filtering SNPs were as follows, minor allele frequency (MAF) , missingness , and biallelic. Three outlier genotypes were found in an initial principal components analysis (PCA) of the *C. maxima* data and were removed, as they were likely not *C. maxima*. Variants were further filtered for specific uses as described below.

## Population Genomics Analysis

ADMIXTURE , which uses a model-based approach to infer ancestral populations () and admixture proportions in a given sample, was used to explore population structure in each dataset. ADMIXTURE does not model linkage disequilibrium (LD); thus, marker sets were further filtered to obtain SNPs in approximate linkage equilibrium using the “–indep-pairwise” option in PLINK with set to 0.1, a window size of 50 SNPs, and a 10 SNP step size . All samples labeled as cultivars or breeding material were removed from the data prior to running ADMIXTURE. These samples were removed to prevent structure created through breeding from appearing as ancestral populations. Ancestral populations were then assigned to cultivars after training on data without the cultivars using the program’s projection feature. Cross-validation was used to determine the best value for each species. Briefly, ADMIXTURE was run with different values (1-20) and the cross-validation error was reported for each . The value with minimal cross-validation error was chosen for each species as shown in Figure [[fig:1]](#fig:1).

Principal components analysis (PCA) was used as a model-free way of determining population structure. PCA was conducted using SNPRelate on the same LD-pruned data used by ADMIXTURE.

Linkage disequilibrium was calculated in each germplasm panel using VCFtools with the settings "–geno-r2 –ld-window 1000". Filtered, but not pruned, data were used for the LD calculation.

## Analysis of Phenotypic Data

Historical data were obtained from the USDA Germplasm Resources Information Network (GRIN; www.ars-grin.gov) for *C. maxima*, *C. pepo*, and *C. moschata*. All duplicated entries were removed for qualitative traits, where categories are mutually exclusive, leaving only samples with unique entries for analysis. Two traits: adult and nymph squash bug damage in *C. pepo* where transformed using the boxcox procedure. Contemporary phenotypic data were collected from a subset of the *C. pepo* collection grown in the summer of 2018 in Ithaca, NY. Field-grown plants were phenotyped for vining bush habit at three different stages during the growing seasons to confirm bush, semi-bush or vining growth habit. Plants that had a bush habit early in the season but started to vine at the end of the season were considered semi-bush.

## GWAS

Variant data were filtered to MAF and missingness , and then imputed prior to association analysis. LinkImpute , as implemented by the TASSEL “LDKNNiImputatioHetV2Plugin” plugin was used for imputation with default settings. Any data still missing after this process were mean imputed. The GENESIS R package, which can model both binary and continuous traits, was used for conducting the associations. All models included the first two PCs of the marker matrix as fixed effects and modeled genotype effect () as a random effect distributed according to the kinship () matrix ()). Binary traits were modeled using the logistic regression feature of GENESIS. The kinship matrix was calculated using A.mat from rrBLUP with mean imputation.

## Genomic Heritability

An estimate of genomic heritability () was calculated for all ordinal and quantitative traits using an equivalent model to what was used for GWAS, but without fixed effects. Variance components from the random genetic effect () and error () were then used to calculate the heritability as .

## Syntenty of Bu putative region in C. pepo and C. maxima

A candidate gene for dwarfism, Bu, in *C. maxima* was elucidated by a previous study and was named Cma\_004516. The Cucurbit Genomics Database gene ID of Cma\_004516 was identified by using the BLAST tool to align primer sequences used for RT-QPCR in the previous study against the *C. maxima* reference genome. The synteny analysis was done by using the Synteny Viewer tool and evaluating *C. maxima*’s chromosome 3 with *C. pepo*’s chromosome 10 and searching for an ortholog to the candidate gene. The physical position of the *C. pepo* ortholog was identified by searching the gene using the Search tool. All tools used in the analysis can be found on the Cucurbit Genomics Database at [cucurbitgenomics.org](https://www.cucurbitgenomics.org).

## Identification of a Core Collection

Subsets representative of each panel’s genetic diversity were identified using GenoCore with the filtered SNP sets. The GenoCore settings were “-cv 99 -d 0.001”.

# Results

## Genotyping

Each *Cucurbita* ssp. collection was genotyped using the GBS approach. The collections comprised 534 accessions for *C. maxima*, 314 for *C. moschata*, and 829 for *C. pepo*. Figure [[fig:2]](#fig:2) shows the geographical distribution of accessions broken down by species. *C. maxima* and *C. moschata* constitute the majority of accessions collected from Central and South America, whereas *C. pepo* accessions are more prevalent in North America and Europe. *C. pepo* had the highest number of raw SNPs (88,437) followed by *C. moschata* (72,025) and *C. maxima* (56,598). After filtering, *C. pepo* and *C. moschata* had a similar number of SNPs, around 30,000, whereas *C. maxima* had an order of magnitude fewer filtered SNPs (1599). This discrepancy may be an artifact of using PstI, a rarer base-cutter previously optimized for GBS of *C. maxima* , rather than ApeKI which was used for *C. pepo* and *C. moschata*. The number and distribution of SNPs across each chromosomes is shown in Table [[tbl:1]](#tbl:1). Maps of SNP distribution for each species are shown in Figure [[fig:3]](#fig:3)

## Population Structure and Genetic Diversity

Filtered SNPs were used for population structure analysis. Available geographical, phenotypic, and other metadata were retrieved from GRIN and were used to help interpret structure results. Results from model-based admixture analysis are shown in Figure [[fig:4]](#fig:4) panel a. These data support 10 ancestral groups (K=10) in C. pepo, 6 in *C. moschata*, and 6 in *C. maxima*. Population structure was driven mostly by geography, except in *C. pepo* where the presence of different subspecies was responsible for some of the structure. Commonalities among structure groups are described in Table [[tbl:2]](#tbl:2). The first two principal components (PCs) of the marker data are shown in Figure [[fig:4]](#fig:4) panel b. As with the model-based analysis, PCA showed geography as a main driver of population structure with accessions being derived from Africa, the Arab States, Asia, Europe, North America, and South/Latin America. PC1 in *C. pepo* separates *C. pepo* subsp. *ovifera*, which have a North American Origin, from subsp. *pepo*.

Ancestry proportions from admixture analysis were projected onto cultivars/market types identified in the accessions. Cultivars were grouped according to known market class within species to help identify patterns in ancestry among and between market classes. Key market types identified in accessions from *C. pepo* include Acorn, Scallop, Crook, Pumpkin (Jack O’ Lantern), Zucchini, Marrow, Gem, and Spaghetti; Neck, Cheese, Japonica, and Calabaza in *C. moschata*; and Buttercup, Kobocha, Hubbard, and Show (Giant squash) in *C. maxima*. These groupings are shown in Figure [[fig:5]](#fig:5). In general, members of each market class exhibit similar ancestry proportions. In *C. pepo*, market classes from the two different subspecies had distinct ancestry patterns. For example, Acorn, Scallop and Crook market classes are all from subsp. *ovifera* and all of these classes had similar ancestry proportions with roughly 20% of ancestry from the wild *ovifera*. In contrast, market classes within *pepo* had a small percentage of ancestry from wild *ovifera* and more ancestry in common with European and Asian accessions. With *C. moschata*, Neck, Cheese, and Calabaza market classes showed very similar ancestry patterns, whereas the Japonica class was more distinct. Relative to the *C. pepo* and *C. moschata*, the *C. maxima* cultivars were less differentiated from one another.

Results from linkage-disequilibrium analysis are shown in Figure [[fig:6]](#fig:6). Similar trends are seen across species. In general, LD decays to zero once the distance between markers reaches more than 2 megabases (Mb). *C. pepo* maintains a higher LD, with an average R-squared between markers of 0.1 even beyond 2 Mb.

## Analysis of Phenotypic Data

All historical phenotypic data from GRIN were compiled for analysis. Only traits with 100 entries were considered for further analysis. Filtering resulted in 26 traits for *C. pepo*, 5 for *C. moschata* and 16 for *C. maxima*. Traits spanned fruit and agronomic-related characteristics, as well as pest resistances. The number of records for a given trait ranged from 108 to 822, with an average of 270. Fruit traits included fruit width, length, surface color and texture, and flesh color and thickness. Agronomic data included plant vigor and vining habit, and several phenotypes related to maturity. Pest-related traits included susceptibility to cucumber beetle and squash bug in *C. pepo* and Watermelon mosaic virus (WMV) and powdery mildew (PM) in *C. maxima*. Figure [[fig:7]](#fig:7) shows the distribution for each quantitative trait.

## Genome-wide Association

Genome-wide association studies were conducted for all traits using a standard mixed-model K + Q analysis. A weak signal was detected in *C. moschata* on chromosome 3 for fruit length. Weak signals were detected in *C. maxima* for fruit ribbing on chromosome 17 and green fruit on chromosome 20. Five phenotypes were significantly associated with SNPs in *C. pepo*: bush/vine plant architecture on chromosome 10 using contemporary and historic data, fruit flesh thickness on chromosome 2, green fruit on chromosomes 2 and 19, and a non-significant, but clear signal for flesh color on chromosome 5. The bush/vine phenotype exhibited the strongest signal. Manhattan plots for the GWAS results are shown in Figure [[fig:8]](#fig:8), and the corresponding quantile-quantile plots are shown in Figure [[fig:9]](#fig:9).

## Genomic Heritability

An estimate of genomic heritability was calculated for all quantitative and ordinal traits. In *C. pepo*, seed weight and morphological traits such as fruit length and width had very high () heritability estimates. Disease and insect resistance traits had lower heritabilites from 0.181-0.228. Trends were similar in both *C. moschata* and *C. maxima*, with *C. maxima* having lower heritability estimates across the board.

## Syntenty of Bu putative region in *C. pepo* and C. *maxima*

A candidate gene for dwarfism found in the species *C. maxima* was named Cma\_00451623 and corresponds to the gene ID CmaCh03G013600 in the Cucurbit Genomics Database. The gene Cp4.1LG10g05740 on chromosome 10 in *C. pepo* was found to be orthologous to CmaCh03G013600 and coincides with the region significantly associated with the bush/vine plant architecture phenotype identified by GWAS in the *C. pepo* collection.

## Development of a Core Collection

A core set of accessions that covered over 99% of total genetic diversity was identified in each of the panels. Roughly 5%-10% of the accessions were required to capture the genetic diversity in the panels (see Figure [[fig:10]](#fig:10). This amounted to 117 accessions in *C. pepo*, 72 in *C. moschata*, and 72 in *C. maxima*.

# Discussion

*Cucurbita pepo*, *C. moschata*, and *C. maxima* exhibit a wide range of phenotypic diversity. This diversity was evident in the GRIN phenotypic records for these species. We have demonstrated that there is also a wide range of genetic diversity through genotyping-by-sequencing and genetic analysis of available specimens from the germplasm collections. Thousands to tens of thousands of whole-genome markers where discovered for each species. Clustering of samples and admixture analysis produced results that align closely with known secondary centers of origin in all species. This was especially clear in our analysis of the *C. pepo* collection. *Cucurbita pepo* has its origin in the new world, with a secondary center of diversification in Europe. This pattern was conspicuous in the our PCA analysis. Analysis of the admixture patterns within common market classes mirrored the results of the broader diversity panel. For example, it is well known that the Acorn, Scallop and Crook type *C. pepo* were primarily developed in the Americas, whereas Zucchini, Marrow, and Gem squash were developed in Europe. Thus, it is not surprising that Acorn, Scallop, and Crook types have a large proportion of subsp. *ovifera* in their background. Likewise, the Neck, Cheese, and Calabaza types have their origins in the Americas, whereas the Japonica type has more shared ancestry with Asian landraces. The various *C. maxima* market classes were less distinct from one another. Morphologically, many of the classes (Buttercup, Kabocha, and Kuri) are very similar, so it is not surprising that their admixture proportions are similar.

Linkage decay curves showed a common pattern across all species, with the correlation between markers falling off precipitously around 2 Mb. Relative to the other two species, *C. pepo* had a higher baseline LD. This is likely due to the presence of two distinct subspecies, subsp. *ovifera* and susbsp. *pepo*, in the *C. pepo* panel. In general, the three *Cucurbita* species studied have much higher LD than other outcrosses, such as maize. Studies in maize have shown that LD drops off within kilobases rather than megabases in diverse accessions . This suggests that the effective population size of *Cucurbita* species is much smaller than other agricultural species. Further, although we have fewer markers in *C. maxima*, it is likely that the number of markers is sufficient to pick up major population structure in the panel given the extent of LD.

Our GWAS analysis using contemporary and historic plant habit data led to the mapping of a locus on chromosome 10 associated with the bush/vine phenotype. This locus is likely the bush gene (Bu) locus that has been finely mapped to this location in previous *C. pepo* studies . Although our GWAS hit does not a constitute a novel gene association, it does demonstrate that the Bu locus, previously mapped in biparental populations, is also the primary driver of the bush phenotype in diverse germplasm. Thus, this locus is likely to have utility across a wide array of germplasm. We also demonstrated that this locus is syntenic with the bush gene previously mapped in *C. maxima* .

Although few signals were detected for traits outside of plant height, this is not a indication that there is an issue with the phenotypic data. Instead, these results may indicate that many of the traits either have a more quantitative nature or are confounded with population structure in some way that cannot be corrected. Our genomic heritability estimates were high for many of the morphological and agronomic traits in each species; yet, no major loci were detected via GWAS. This suggests that many of these traits have a more complex trait architecture.

Our data provides many genome-wide markers which could be used to develop marker panels for use in breeding applications, as has been done in other crops . Possible breeding applications would include marker assisted selection, marker assisted backcrossing, and purity assessment of seedstock using a low density panel; whereas, a medium density panel could be developed for routine genomic selection . Our clustering of samples based on marker data suggest geography is a key driver for overall population structure. When projecting ancestry proportions onto cultivars of known market classes, the ancestry proportions were relatively similar within market class grouping. Although there is genetic diversity within each species, this diversity is constrained within market classes. This suggests that crosses between these market classes would greatly increase the amount of genetic diversity to be leveraged in breeding efforts. Crossing between market classes would come at the cost of bringing in undesirable characteristics with regard to achieving a specific morphotype associated market class. This cost could be mitigated through the use of markers to recover morphotype expeditiously during pre-breeding .

Our data provides a useful starting point for future studies. In the case where traits are common in the panel, the panel can be phenotyped for a trait of interest and combined with marker data and insight provided by our study. We demonstrated this approach in our association analysis of the bush gene. In the case of a rare phenotype, such as a resistance gene, subsets of the germplasm and markers should be used to develop custom populations. Plant introductions (PI) are frequently used as source parents in mapping studies and for germplasm improvement, as was the case for mapping Phytophthora capsici resistance and developing resistant breeding lines . We found some traits that had high heritability, such as morphological traits, but we were not able to find any associations. This is consistent with other studies looking at morphological traits in squash . Genomic predication rather than association may be the best approach for these traits. Certain applications, such as creation of a hapmap or diversity atlas, require higher density re-sequencing data. Our GenoCore analysis provides a subset of several hundred accessions that will be informative for re-sequencing efforts.

# Conflict of Interest Statement

Michael Mazourek is a co-founder of Row 7 Seeds, but neither receives compensation nor holds equity.

# Author Contributions

COH wrote the first draft. MM, ZF, and RG provided project oversight. COH, JF, and KB conducted data analysis. KR, JL, and BJ assisted with data curation and germplasm selection. All authors contributed to the article.

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# Data Availability Statement

The datasets generated for this study includeing variant and raw sequence data are available on the Cucurbit Genomics Dataase at [cucurbitgenomics.org](https://cucurbitgenomics.org). The phenotypic data used are available for download from the USDA Germplasm Resources Information Network (GRIN; [www.ars-grin.gov](https://www.ars-grin.gov)). Intermediate files and code used in the study are available on Github at [www.github.com/ch728/Cucurbita-USDA](https://www.github.com/ch728/Cucurbita-USDA).

# Figure captions

# Tables

Distribution and number of raw and filtered SNPs per chromosome for each species [tbl:1]

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| Chromosome | Raw | Filtered | Raw | Filtered | Raw | Filtered |
| 0 | 12498 | 2550 | 2708 | 546 | 1501 | 132 |
| 1 | 7497 | 2831 | 3890 | 1468 | 4185 | 121 |
| 2 | 5153 | 2049 | 3661 | 1538 | 2101 | 55 |
| 3 | 4875 | 1943 | 3472 | 1499 | 2201 | 51 |
| 4 | 4598 | 1982 | 6880 | 2553 | 5703 | 106 |
| 5 | 4045 | 1628 | 2716 | 887 | 3115 | 46 |
| 6 | 3871 | 1384 | 3262 | 1159 | 3035 | 92 |
| 7 | 3129 | 1222 | 2668 | 969 | 2705 | 62 |
| 8 | 3875 | 1583 | 2348 | 810 | 2391 | 61 |
| 9 | 3766 | 1390 | 3106 | 995 | 2750 | 84 |
| 10 | 3585 | 1488 | 3550 | 1327 | 2297 | 52 |
| 11 | 3227 | 1216 | 4336 | 1830 | 3713 | 131 |
| 12 | 3089 | 1163 | 3711 | 1330 | 2026 | 47 |
| 13 | 3434 | 1350 | 3106 | 1280 | 2131 | 82 |
| 14 | 3543 | 1291 | 4753 | 1929 | 4317 | 100 |
| 15 | 2640 | 960 | 3564 | 1321 | 2662 | 58 |
| 16 | 3088 | 1060 | 2933 | 1107 | 2058 | 100 |
| 17 | 2994 | 1175 | 2885 | 1096 | 2195 | 86 |
| 18 | 3053 | 1258 | 3341 | 1316 | 1826 | 46 |
| 19 | 3381 | 1340 | 2638 | 990 | 1793 | 46 |
| 20 | 3096 | 1155 | 2497 | 903 | 1893 | 41 |
| Total | 88437 | 32018 | 72025 | 26853 | 56598 | 1599 |

Commonalities among accessions in each group, most groupings are dictated by geography [tbl:2]

|  |  |  |  |
| --- | --- | --- | --- |
|  | *C. pepo* | *C. moschata* | *C. maxima* |
| 1 | Mixed Group; Many from Spain, Turkey, and Syria | Mostly from Mexico | Mixed; Primarily from South America and Asia |
| 2 | Wild subsp. *ovifera* var. *texana* and var. *ozarkana*; North American | Mostly Mexico and Guatemala | Mixed; Primarily from Asia and Europe |
| 3 | Majority from Turkey | Mostly from Mexico | Mostly from North Macedonia |
| 4 | Majority from North Macedonia | Mostly from Africa | Mostly from Argentina |
| 5 | Majority from Egypt | Mostly from India | Mostly Turkey, Iran, Afghanistan |
| 6 | Majority from Mexico | Mixed origin Europe and Americas;Many similar to cheese or neck type | Mostly from Africa |
| 7 | Majority from Syria |  |  |
| 8 | Majority from Pakistan and Afghanistan |  |  |
| 9 | Majority from Spain |  |  |
| 10 | Wild subsp. *fraterna*; Central American |  |  |

Commonalities among accessions in each group, most groupings are dictated by geography [tbl:3]

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Species** | **Trait** | **Type** | **Description** |  |
| *C. pepo* |  |  |  |  |
|  | seed\_wt | Quantitative | Weight of 100 seeds in grams | 0.95 |
|  | plant\_type | Binary | Historical plant architecture data coded as vining or bush | NA |
|  | plant\_type2 | Binary | Contemporary plant architecture data coded as vining or bush | NA |
|  | max\_vig | Ordinal | Maximum plant vigor on 1-5 scale | 0.588 |
|  | min\_vig | Ordinal | Minimum plant vigor on 1-5 scale | 0.618 |
|  | max\_width | Quantitative | Maximum fruit width in centimeters | 0.937 |
|  | width\_min | Quantitative | Minimum fruit width in centimeters | 1 |
|  | len\_max | Quantitative | Maximum fruit length in centimeters | 0.748 |
|  | len\_min | Quantitative | Minimum fruit length in centimeters | 0.841 |
|  | flesh\_max | Ordinal | Maximum fruit thickness in centimeters | 0.614 |
|  | flesh\_min | Ordinal | Minimum fruit thickness in centimeters | 0.425 |
|  | sb\_nymph | Quantitative | Number of squash bug nymphs on plant | 0.181 |
|  | sb\_adult | Quantitative | Number of adult squash bugs on plant | 0.206 |
|  | cuc\_inj | Ordinal | Severity of beetle damage on a 0-4 scale | 0.228 |
|  | or\_flesh | Binary | Flesh color coded as orange or not orange | NA |
|  | yl\_flesh | Binary | Flesh color coded as yellow or not yellow | NA |
|  | yl\_fruit | Binary | Color of fruit coded as yellow or not yellow | NA |
|  | tan\_fruit | Binary | Color of fruit coded as tan or not tan | NA |
|  | gn\_fruit | Binary | Color of fruit coded as green or not green | NA |
|  | globe\_fruit | Binary | Fruit shape as globe or not globe | NA |
|  | oblong\_fruit | Binary | Fruit shape as oblong or not oblong | NA |
|  | smooth\_fruit | Binary | Furit texture as smooth or not smooth | NA |
|  | rib\_fruit | Binary | Degree of ribbing | NA |
|  | spec\_fruit | Binary | Fruit patterning as speckled or not speckled | NA |
|  | mot\_fruit | Binary | Fuit patterning as mottled or not mottled | NA |
|  | solid\_fruit | Binary | Fruit patterning as solid color or patterned | NA |
| *C. moschata* |  |  |  |  |
|  | fruit\_len | Quantitative | Fruit length in centimeters | 0.804 |
|  | fruit\_diam | Quantitative | Fruit diameter in centimeters | 0.478 |
|  | maturity | Binary | Fruit maturity as early or late | NA |
|  | or\_fruit | Binary | Fruit color coded as orange or not orange | NA |
|  | smooth\_fruit | Binary | Fruit surface texture encoded as smooth or not smooth | NA |
| *C. maxima* |  |  |  |  |
|  | len | Quantitative | Fruit length in centimeters | 0.374 |
|  | set | Ordinal | Fruit set from poor to excellent (1-9) | 0.254 |
|  | diam | Quantitative | Fruit diameter in centimeters | 0.249 |
|  | watermelon\_mosaic | Ordinal | Susceptibility to WMV from slight to severe (0-9) | 0.18 |
|  | cuc\_mosaic | Ordinal | Cucumber mosaic susceptibility from slight to severe (0-9) | 0.129 |
|  | maturity | Quantitative | Number of days from field transplanting to date of first pollination | 0.388 |
|  | unif | Ordinal | Fruit uniformity from poor to excellent (1-9) | 0.157 |
|  | pm | Ordinal | Susceptibility to PM from slight to severe (0-9) | 0.192 |
|  | plant\_habit | Binary | Plant type as vining or not vining | NA |
|  | vig | Ordinal | Plant vigor from poor to excellent (1-9) | NA |
|  | or\_flesh | Binary | Flesh color as orange or not orange | NA |
|  | rib | Ordinal | Fruit ribbing from slight to pronounced (1-9) | 0.427 |
|  | fruit\_spot | Ordinal | Fruit spotting from slight to pronounced (1-9) | 0.132 |
|  | gray\_fruit | Binary | Fruit color encoded as gray or not gray | NA |
|  | or\_fruit | Binary | Fruit color encoded as orange or not orange | NA |
|  | gn\_fruit | Binary | Fruit color encoded as green or not green | NA |
|  |  |  |  |  |