Genetic Diversity and Population Structure of Cucumber (Cucumis sativus L.)

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Abstract

Knowing the extent and structure of genetic variation in germplasm collections is essential for the conservation and utilization of biodiversity in cultivated plants. Cucumber is the fourth most important vegetable crop worldwide and is a model system for other Cucurbitaceae, a family that also includes melon, watermelon, pumpkin and squash. Previous isozyme studies revealed a low genetic diversity in cucumber, but detailed insights into the crop’s genetic structure and diversity are largely missing. We have fingerprinted 3,342 accessions from the Chinese, Dutch and U.S. cucumber collections with 23 highly polymorphic Simple Sequence Repeat (SSR) markers evenly distributed in the genome. The data reveal three distinct populations, largely corresponding to three geographic regions. Population 1 corresponds to germplasm from China, except for the unique semi-wild landraces found in Xishuangbanna in Southwest China and East Asia; population 2 to Europe, America, and Central and West Asia; and population 3 to India and Xishuangbanna. Admixtures were also detected, reflecting hybridization and migration events between the populations. The genetic background of the Indian germplasm is heterogeneous, indicating that the Indian cucumbers maintain a large proportion of the genetic diversity and that only a small fraction was introduced to other parts of the world. Subsequently, we defined a core collection consisting of 115 accessions and capturing over 77% of the SSR alleles. Insight into the genetic structure of cucumber will help developing appropriate conservation strategies and provides a basis for population-level genome sequencing in cucumber.

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Introduction

Commonly known as cucubits or gourds, the botanical family Cucurbitaceae includes a number of cultivated species of global or local economical importance [1]. Cucumber (Cucumis sativus L.) is the fourth most important vegetable worldwide [2]. As the first in cucubits, the cucumber genome sequence elucidated chromosomal evolution in the genus Cucumis and afforded novel insights into several important biological processes such as biosynthesis of cucurbitacin and “fresh green” odor. Cucumber is being developed as a new model species in plant biology due to its small number of genes, rich diversity of sex expression, suitability for vascular biology studies, short life cycle (three months from seed to seed), and accumulating resources in genetics [3,4] and genomics [5,6].

The rapid advance of Next Generation Sequencing (NGS) technologies makes it affordable to re-sequence multiple genotypes of a given species to generate a haplotype map that displays the genome-wide patterns of genetic variation at a single base resolution [7,8,9]. However, such population sequencing requires...
a well-designed sampling of the species' genetic diversity. To date, little is known about the genetic diversity and structure in cucumber.

Cucumber is indigenous to India [10] and likely originated from the foothills of the Himalayan Mountain, where its only two botanical varieties both were discovered, namely the domesticated cucumber C. s. var. sativa and the wild cucumber C. s. var. hardwickii (Royce) Alef. The cultivation of cucumber seems to have spread rapidly from India to Western Asia, and then to Southern Europe. It has been reported that cucumber was introduced into China through the Silk Route by the diplomat Zhang Qian (164 B.C.–114 B.C.) during the Han Dynasty and subsequently spread to East Asia [11]. A unique semi-wild landrace, named Xishuangbanna Gourd, was discovered in the Prefecture Xishuangbanna of the Province Yunnan in Southwest China. To date, its relationship to other phylogeographic populations is unknown.

Genetic diversity in the U.S. cucumber collection (~1,000 Plant Introductions, or Pls) maintained by the National Plant Germplasm System (NPGS) was investigated in a series of studies by analysis of variation at 21 isozyme loci [12,13,14,15]. Overall, these studies revealed low diversity (~2.2 alleles per polymorphic locus) in domesticated cucumber. Possibly due to the lack of polymorphism, the isozyme data did not allow consistent clustering of the Pls. Consequently, the genetic structure of domesticated cucumber remained largely unknown.

In this study we assembled a mega-collection consisting of all available cucumber accessions from the national germplasm centres of China, the Netherlands and U.S. We fingerprinted a total of 3,342 accessions with 23 highly polymorphic Simple Sequence Repeat (SSR) markers. The SSR data were used to assess overall genetic diversity and to reveal a clear population structure in cucumber. In addition, the data allowed the construction of a core set of 115 accessions capturing over 77% of the total allelic diversity observed.

Materials and Methods

Materials

A mega-collection of 3,342 cucumber accessions was used in this study. A total of 1,692 accessions were collected from the China National Vegetable Germplasm Bank located at the Institute of Vegetables and Flowers at the Chinese Academy of Agricultural Sciences (IVF-CAAS), 883 accessions from the U.S. National Plant Germplasm System (NPGS, http://www.ars-grin.gov/npgs/), 759 accessions from the Centre for Genetic Resources, the Netherlands (CGN, http://www.cgn.wur.nl), 2 accessions from the IPK (http://www.ipk-gatersleben.de/), and 6 accessions from the East-West Seed International Ltd. The mega-collection included 235 accessions originating from India, 1,361 from China (excluding Xishuangbanna), 208 from Xishuangbanna, 209 from Southeast Asia, 141 from Western Asia, 802 from Europe, 176 from the Americas, 50 from Africa and 8 from Oceania. Additionally, 152 accessions of unknown origin were included in the study. Passport information about the mega-collection is listed in Table S1.

SSR analyses

DNA extraction. For most of the study accessions multiple self-crossing was performed during gene bank curation, and previous studies on cucumber germplasm have revealed limited intra-accession variation. To increase the efficiency in analysing the large number of accessions, leaves from five randomly chosen plants per accession were bulked into a single sample. Genomic DNA was isolated from about one gram of leaf tissue from each bulk using a CTAB-based method [16].

SSR markers. In previous studies [3,4], we determined the polymorphism level of 995 SSR markers using a selected panel of 11 cucumber genotypes that represent six market types worldwide: ‘Chinese Long’, Southern China type, Xishuangbanna type, European greenhouse type, American slicing type, and Japanese type. For the present study, we selected 23 highly polymorphic SSR markers that are evenly distributed across the cucumber genetic map (Table 1). Each selected SSR marker is able to produce a clear, stable, and single polymerase chain reaction (PCR) product. On each chromosome arm, there is at least one SSR (Figure 1). The genetic distance between two neighbouring markers ranges from 7.1 cM to 53.7 cM with an average of 29.4 cM.

PCR amplification. The 23 SSR markers were analyzed using four sets of multiplex PCR reactions. Each multiplex was carefully assembled according to the compatibility of the SSRs during PCR and the molecular size of their amplicons. The forward primer of the SSR markers was labelled with one of the four fluorescent dyes, carboxy fluorescein (FAM), carboxytetramethylrhodamine (TAMRA), hexachloro-6-carboxyfluorescein (HEX) and ROX (carboxy-X-rhodamine). Multiplex PCRs were performed in a 11 µl volume containing approximately 20–50 ng template DNA, 0.4 pmol of each primer, and 5 µl Multiplex PCR Master Mix (QIAGEN Multiplex PCR Kit, Qiagen). Reactions were performed in an ABI 9700 thermocycler with an initial denaturation step of 15 min at 95°C, followed by 35 cycles of 95°C for 30 s, 57°C for 90 s and 72°C for 70 s, and a final extension at 60°C for 30 min. For PCR fragment size determinations, 0.25 µl of an internal size standard (Liz-500, LIZ) was mixed with 1 µl of diluted PCR product (1/100) and 9 µl formamide. The mixture was heated at 94°C for 3 minutes and then cooled within icy water. Electrophoresis was carried out on an ABI 3130xl genetic analyzer (Applied Biosystems). Analyses were performed using the Genemapper 4.0 software.

Data analysis

Genetic structure. The model-based program STRUCTURE [17] was used to infer population structure and to identify hybrid forms. In order to identify the number of populations (K) capturing the major structure in the data, we used a burn-in period of 10,000 Markov Chain Monte Carlo iterations and 100,000 run length, an admixture model following Hardy-Weinberg equilibrium and correlated allele frequencies as well as independent loci for each run. Three independent runs were performed for each simulated value of K, ranging from 1 to 20. Subsequently, an Pr[X|K] index with respect to each K was used to calculate AK using the formula described by Evanno et al. [18]. The optimal K depends on the first peak of AAK = [L(K)/K]/s[Pr(x|k)], where [L(K)/K] denotes the absolute value of the second order rate of change of Pr(x|K), and s[Pr(x|k)] the standard deviation of the Pr(x|K).

Genetic diversity assessment. For the entire collection as well as the subpopulations, allele richness, gene diversity and the number of unique alleles were calculated with PowerMarker version 3.25 [19], using genetic distances calculated by the CS Chord 1967 method [20]. Tree topologies were constructed based on the neighbour-joining method with MEGA [21]. For each subpopulation inferred from STRUCTURE, the observed and effective number of alleles, the observed and expected heterozygosity, as well as Shannon’s information index were calculated by POPGENE. As a measure of genetic differentiation between pairs of sub-populations, we computed FST statistics with the AMOVA
means in Arlequin 3.11 [22]. Principal Component Analysis (PCA) in NTSYSpc2.11 was used to analyse genetic relationships among accessions and to determine the optimal number of clusters in the study.

**Core collection sampling.** For sampling core collections, we used the Maximization (M) algorithm implemented in MSTRAT software version 4.1 [23]. For evaluation of core collection’s minimal size and for individual sampling of the collections, 20 replicates of 20 iterations for each replicate were performed.

**Results**

**Molecular fingerprinting of the mega-collection**

We fingerprinted 3,342 cucumber accessions using 23 SSR markers. In total, 94.9% of PCR reactions yielded clear peaks in electrophoresis and generated 72,960 data points. The remaining 5.1% resulted in multiple bands with low quality and consequently were treated as missing values in subsequent analyses. Out of the 72,960 data points, 61,976 (84.9%) corresponded to single peaks that were regarded as homozygous for the given SSR locus, while 8,929 (12.2%) corresponded to double peaks that were regarded as heterozygous. Only 2,055 (2.8%) of the data points corresponded to more than two clear peaks, of which only the two highest peaks were recorded that were regarded as heterozygous. This simplified scoring may cause under-estimation of rare alleles.

The 23 SSR markers generated a total of 316 alleles in the mega-collection (Table 1). Out of the 316 SSR alleles, 64 displayed a frequency of more than 5% in the total sample and hence were classified as ‘common’ alleles, while another 64 displayed frequencies between 1% and 5% that were denoted as ‘less common’ alleles. Out of the 188 remaining alleles, 121 were denoted as ‘rare alleles’ with frequencies between 0.1% and 1% and 67 as ‘very rare’ alleles with frequencies smaller than 0.1%. That 59% of the observed alleles showed a frequency of less than 1% in the mega-collection suggested that this collection represents the bio-diversity of cucumber broadly.

**Genetic structure in cucumber**

The population structure of the cucumber mega-collection was firstly inferred using STRUCTURE 2.3.1 [18]. The Evanno et al. (2005) correction showing only peak of ΔK, for K = 3, suggested the presence of three main populations in cucumbers (Figure 2A). The classification of accessions into populations by the model-based method was shown in Figure 2B and Table S1. In total 2,932 accessions (87.7%) were assigned to one of the three populations, where more than 70% of their inferred ancestry was derived from one of the model-based populations. Population 1, 2, and 3 (P1, P2, and P3) consisted of 1399, 1129, and 404 accessions, respectively. The remaining 410 accessions (12.3%) were categorized as having admixed ancestry, including 152 admixtures between P1 and P2 (P1P2), 163 between P1 and P3 (P1P3), and 95 between P2 and P3 (P2P3).

The unrooted neighbour-joining tree (Figure 2C) is consistent with the aforementioned model-based population structure. Accessions of the three populations are largely, but not completely, separated. Admixtures are mostly located in between two populations.

We also performed Principal Component Analysis (PCA) on 1,026 accessions without any missing SSR data (Figure 2D). This analysis largely supported the separation of the accessions into three populations. P3 can be divided into two subgroups, one consisting mainly of accessions from India and the other with accessions from Xishuangbanna. Most admixtures appear in between the two populations.

In summary, the model-based ancestry analysis, the phylogenetic tree and the PCA strongly supported that cucumber has three well-differentiated genetic populations and admixtures.

**Geographical distribution of cucumber populations**

The classification of populations appeared highly correlated with the geographical distribution of the cucumber accessions (Table 2, Figure 3). P1 includes cucumber germplasm from North and East China, Japan and South Korea, P2 includes accessions...
| Locus  | cM  | Motif | Primer sequence (5’ to 3’) | Na | Ne | Ho  | He  | I  | PIC |
|--------|-----|-------|--------------------------|----|----|-----|-----|----|-----|
| Chr1   | SSR0018 | 19 (AAG) | F:GGGTCTAATATTTGGGATGG  | 16 | 3  | 0.29 | 0.60 | 1.3 | 0.65 |
|        |       |       | R:GGTTGTTCTGTGGAATGTTG   |    |    |      |      |     |     |
| SSR11340 | 48 (AG) | F:TTGTTTTTGTTGGGACTGCA  | 14 | 2  | 0.10 | 0.54 | 1   | 0.76 |
|        |       |       | R:GTGTACACTCCACTCCCTTC   |    |    |      |      |     |     |
| SSR05723 | 78 (AT) | F:GGGTGAATTGGCTTTTCTG  | 10 | 2  | 0.12 | 0.51 | 1.1 | 0.38 |
|        |       |       | R:GGTTCAATACGAGGATGTGC   |    |    |      |      |     |     |
| SSR16695 | 86 (CAT) | F:GGAATGAAACACATCCCAACG | 7  | 2  | 0.13 | 0.52 | 1.2 | 0.39 |
|        |       |       | R:GGTTGTTCTGCTAAGGATGTCC |      |     |      |      |     |     |
| Chr2   | SSR16226 | 20 (TCTT) | F:GGGTAAATTCCCAAACGG   | 8  | 3  | 0.23 | 0.67 | 1.2 | 0.62 |
|        |       |       | R:GGAGAAATTTGAATTCGGCAG  |    |    |      |      |     |     |
| SSR23220 | 36 (AGAAGG) | F:GGGTGGAATCTGGGTTTG  | 10 | 2  | 0.07 | 0.59 | 0.9 | 0.5  |
|        |       |       | R:GGTGGAAATATGGAGGGAGAG  |    |    |      |      |     |     |
| SSR22653 | 73 (AG) | F:TGAATCTCTTTGGTGATCAAA | 14 | 3  | 0.23 | 0.61 | 1.2 | 0.55 |
|        |       |       | R:GGGAGAAGGAGGAGAGTGTG   |    |    |      |      |     |     |
| SSR23370 | 95 (TA) | F:GATTATGAGGATGAACCACC   | 13 | 3  | 0.31 | 0.70 | 1.5 | 0.56 |
|        |       |       | R:GCCAAACACTTCTCTCTATGCAAC |      |     |      |      |     |     |
| Chr3   | SSR01738 | 2.2 (GAA) | F:GGGTAGAAGAATACCAATAGG  | 13 | 2  | 0.15 | 0.50 | 0.9 | 0.42 |
|        |       |       | R:GGCTAAGAATATGATAGAACC  |    |    |      |      |     |     |
| SSR16056 | 56 (CACCCCT) | F:GGGTGGAATCTGGGTTTG  | 15 | 4  | 0.29 | 0.74 | 1.6 | 0.68 |
|        |       |       | R:GGTGGAAATATGGAGGGAGAG  |    |    |      |      |     |     |
| SSR05012 | 105 (TCT) | F:TTTAAGGGCTGCAATAAAGTGT | 6  | 2  | 0.36 | 0.51 | 0.7 | 0.34 |
|        |       |       | R:GGTTCCAAATTAAAGCTTCCC  |    |    |      |      |     |     |
| Chr4   | SSR11043 | 4.4 (GAA) | F:TACACTCTTGGAGAGGACC   | 15 | 3  | 0.33 | 0.66 | 1.5 | 0.64 |
|        |       |       | R:GGTTACACTCTCTTTACCGG  |    |    |      |      |     |     |
| SSR05125 | 25 (CACCCCT) | F:GGGTGGAATCTGGGTTTG  | 14 | 2  | 0.12 | 0.58 | 1.1 | 0.68 |
|        |       |       | R:GGTGGAAATATGGAGGGAGAG  |    |    |      |      |     |     |
| SSR07543 | 35 (GA) | F:GGGTGGTTCTCCTCTCACTC  | 9  | 3  | 0.16 | 0.63 | 1.1 | 0.63 |
|        |       |       | R:GGTTCCAAAATGAAAATCAGC  |    |    |      |      |     |     |
| Chr5   | SSR19998 | 6 (AATC) | F:CTTTGGCAAGCCTCTCACC  | 10 | 1  | 0.21 | 0.30 | 0.7 | 0.3  |
|        |       |       | R:GGTTTGCTTGGGCTTCCAG  |    |    |      |      |     |     |
| SSR16068 | 13 (AG) | F:GTGCAAAAGGGGAGTGGTTG   | 12 | 2  | 0.14 | 0.50 | 0.9 | 0.57 |
|        |       |       | R:GGTTGCTGTGTTGCTGTGTTG  |    |    |      |      |     |     |
| SSR02895 | 54 (AT) | F:GGAGAAAGGATGTTGCAAGTGC | 23 | 4  | 0.30 | 0.73 | 1.7 | 0.88 |
|        |       |       | R:GGAGGAATGTGGGATGAGAG  |    |    |      |      |     |     |
| Chr6   | SSR20852 | 6.2 (TA) | F:GGTTCCTTTCAGGCGTACG   | 24 | 3  | 0.13 | 0.68 | 1.7 | 0.74 |
|        |       |       | R:GGGCTTCAATTTAGGAACC  |    |    |      |      |     |     |
| SSR31399 | 40 (AT) | F:AGCTCCGAGGATACCCACTCT | 10 | 3  | 0.18 | 0.61 | 1.1 | 0.67 |
|        |       |       | R:AGAAGAAGACCTGGAACAGCA  |    |    |      |      |     |     |
| SSR20218 | 90 (AGA) | F:TGGCCACGTGCTCTATATC  | 8  | 2  | 0.25 | 0.54 | 0.9 | 0.6  |
|        |       |       | R:GCTAAATGGAGGGGAGAGGA  |    |    |      |      |     |     |
| Chr7   | SSR29620 | 5.6 (GAGATG) | F:GTGCTTGGAGGATGCTTTCTG | 11 | 2  | 0.19 | 0.49 | 1   | 0.64 |
|        |       |       | R:GGTTTTATGAGGATGTTTGGT  |    |    |      |      |     |     |
| SSR14861 | 32 (ATAC) | F:CGGTTAGCTCTGGTTGAAATG | 16 | 3  | 0.15 | 0.59 | 1.3 | 0.7  |
|        |       |       | R:GGTAAAGGCGGAGGAAAAACAC |      |     |      |      |     |     |
| SSR13787 | 59 (AT) | F:GCAACTCCAACACATCCCTC  | 25 | 3  | 0.35 | 0.66 | 1.7 | 0.76 |
|        |       |       | R:GGCAGCTAATACCTCTCACC  |    |    |      |      |     |     |

Na: the observed number of alleles; Ne: the effective number of alleles; Ho: observed heterozygosity; He: expected heterozygosity; I: Shannon’s information index; PIC: polymorphism information content.  
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collected from Central and West Asia, Europe, America and Africa, while P3 comprises 404 accessions mainly from India and Xishuangbanna. Xishuangbanna is geographically close to the Indo-Gangetic plain where cucumber originated. Also the distribution of the admixtures appeared related to geography. About a quarter of the P1P2 admixtures (36 of 152) originated from East Europe including the Russian Federation and the Former Soviet Union. The P1P3 admixtures are mainly from Southwest China and India, and the P2P3 admixtures mainly from India. The admixtures are most likely due to natural or artificial hybridization.

In PCA analysis (Figure 2D), P3 can be divided into two subgroups, one with accessions from India mainly and the other with accessions from Xishuangbanna. With more markers, the subgroups might be differentiated into two separate populations.

Genetic diversity assessment

AMOVA analysis of the populations P1, P2 and P3 indicated that 68.1% of the variation was due to differences within populations, while 31.9% was due to differences among populations (Table 3). Pairwise estimates of $F_{ST}$ using AMOVA indicated a high degree of differentiation between the three model-based populations with values ranging from 0.30 to 0.33 (between P1 and P2: 0.33, P1 and P3: 0.30; P2 and P3: 0.31).

Genetic parameters for all accessions and the three model based populations are given in Table 4. For all accessions genotyped, the average number of alleles per locus was 13.70, ranging from 6 (SSR05012) to 25 (SSR13787) (Table 1). The Expected number of alleles was much smaller than that of observed, indicating the existence of few high frequency alleles in cucumber genome. The overall genetic diversity of cucumber was similar to tomato (0.2–
0.82) [26].

58
Among the three model based populations, P3 possessed the highest diversity (He = 0.52), though there was no significant differences in Na and Ne. Even with the least number of accessions involved, P3 displayed the largest number of population specific alleles (n = 40). This supports the notion that cucumber was originated from India [10].

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For all populations, the observed heterozygosity was lower than expected heterozygosity, which may be caused by inbreeding during seed propagation by germplasm banks.

62
Core Collection
64
The Maximization or M strategy algorithm was used to select the core set of the mega-population. The M curve reached the plateau when approximately 115 accessions were selected (Figure 4A). Taking geographic distribution, existing phenotype data, and population structure into account, we selected 115 accessions to define the core collection (Table S2). The core collection captures 77.2% of the observed SSR alleles.

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The dataset allowed us to visualize the population structure of cucumber and to assemble a core collection of 115 accessions that cover over 77% of the observed SSR alleles.

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Out of the 316 SSR alleles scored, 59% and 80% are present in less than 1% and 5% of the mega-collection, respectively. As the level of gene flow is negatively related to the number of rare alleles [27], the abundance of rare alleles in the mega-collection indicates that there has been limited genetic exchange among sub-populations in cucumber. The results also supports that independent migrations of cucumber have occurred from India to other parts of the world. The migration route to Europe and then to the Americas and Africa are clearly through Central and West Asian countries. Cucumber accessions from China and other East Asian countries are genetically differentiated from those in Central and West Asia. It is well documented in Chinese history that cucumber was introduced from there through the Silk Route by the diplomat Zhang Qian (164 B.C.-114 B.C.) [11]. In addition, cucumber cultivation in heated houses during the winter season was recorded in poems from the Tang Dynasty (618–907). These records support that the crop has been intensively cultivated and therefore selected by Chinese farmers for more than one thousand years, which likely caused the genetic differentiation of Chinese (East Asian) cucumbers related to other populations.

73
Cucumber plants in India and Xishuangbanna bear short, ovary to round fruits with sparse spines (Figure S1). The occurrence of black spines, an unfavourable market trait, within Indian resources, is more frequent than the other two populations. Cucumbers should have undergone differential human selection when introduced to other parts of the world. For instance, in Europe and America, short-fruited gherkins and long fruited

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Discussion
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In this study, we carefully selected 23 highly polymorphic SSR markers and used them to fingerprint a large cucumber collection. The dataset allowed us to visualize the population structure of cucumber and to assemble a core collection of 115 accessions that cover over 77% of the observed SSR alleles.

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Geographic origin of the accessions assigned by the software STRUCTURE to populations (P1–P3) and admixtures (P1P2–P2P3).

82
Table 2.

83
| Region          | P1 | P2 | P3 | P1P2 | P1P3 | P2P3 | Total |
|-----------------|----|----|----|------|------|------|-------|
| East Asia       | 1299 | 7  | 4  | 47   | 143  | 3    | 1503  |
| Southeast Asia  | 0  | 8  | 41 | 2    | 5    | 11   | 67    |
| Oceania         | 1  | 4  | 1  | 0    | 1    | 1    | 8     |
| Africa          | 1  | 38 | 4  | 2    | 1    | 4    | 50    |
| America         | 8  | 144| 0  | 22   | 0    | 2    | 176   |
| Europe          | 15 | 716| 3  | 64   | 0    | 4    | 802   |
| Central and West Asia | 1 | 136| 0  | 3    | 0    | 1    | 141   |
| Xishuangbanna   | 1  | 0  | 205| 0    | 2    | 0    | 208   |
| India           | 1  | 16 | 134| 7    | 11   | 66   | 235   |
| Others          | 72 | 59 | 13 | 5    | 0    | 3    | 152   |
| Total           | 1399 | 1129 | 404| 152  | 163  | 95   | 3342  |

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Table 4. Summary of SSR diversity parameters of the model-based populations and admixtures.

88
| Region          | P1 | P2 | P3 | P1P2 | P1P3 | P2P3 | All  |
|-----------------|----|----|----|------|------|------|------|
| Na              | 10.3| 8.65| 10.1| 6.78 | 8.09 | 6.83 | 13.70|
| Ne              | 2.25| 1.79| 2.60| 2.32 | 2.37 | 2.39 | 2.52 |
| Ho              | 0.26| 0.15| 0.14| 0.23 | 0.34 | 0.19 | 0.21 |
| He              | 0.43| 0.37| 0.52| 0.53 | 0.54 | 0.54 | 0.58 |
| I               | 0.90| 0.75| 1.12| 1.01 | 1.06 | 1.07 | 1.18 |
| No. of unique alleles | 17 | 3  | 40 | 0    | 5    | 2    |       |

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Table 3. Analysis of molecular variance (AMOVA) among populations and within populations.

93
| Source of variation | d.f. | Sum of squares | Variance components | Percentage of variation |
|---------------------|------|----------------|---------------------|------------------------|
| Among populations   | 2    | 9543.5         | 2.7                 | 31.9                   |
| Within populations  | 5869 | 33551.8        | 5.7                 | 68.1                   |
| Total               | 5871 | 43095.3        | 8.4                 | 100                    |

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Dutch cucumbers were both developed to satisfy different market need (Figure S1). To enhance total yield, gynoecious varieties were widely adopted by western farmers as they could bear more fruits [28]. But in East Asia where cucumber is often served for direct consumption, characteristics such as proper appearance, crispness and “fresh green” flavour outweigh other traits (Figure S1). As a result, the ‘Chinese Long’ type had been developed to meet with that specific demand.

The Xishuangbanna cucumbers are affiliated to the Indian population. As semi-wild landraces, they apparently lost the unfavourable fruit bitterness but still possess the dependence of short daylight for flowering and fruit setting. The latter trait was discarded in most cucumber cultivars. Accessions from Xishuangbanna are genetically close to each other, suggesting that a single dispersal occurred from India. It is worth mentioning that among the African accessions, six lines are present from Zambia and Zimbabwe classified as P3 and admixture P2P3 (CG7743–CG7745, CG7747–CG7749, Table S1). The accessions were initially misclassified as melon and likely represent another dispersal from India.

The study also has potential implications for cucumber breeding. Currently, most crosses have been made within populations and less frequently among populations. The large differentiation among populations indicates that each population may possess its own alleles and haplotypes. Therefore, crosses between populations will broaden the genetic diversity within current breeding programs and may result in new types of heterosis. This study provides a sound basis for further characterization of the biodiversity of this important vegetable. For example, we are currently re-sequencing the core collection, enabling in-depth discovery of the genetic variation in cucumber.

Table 5. Genetic diversity parameters of the entire sample and those of the core collection based on the 23 investigated SSR loci.

|                | Na  | Ne  | I   | He  |
|----------------|-----|-----|-----|-----|
| All accessions | 13.70 | 2.52 | 1.18 | 0.58 |
| Core collection | 10.65 | 3.24 | 1.44 | 0.66 |

Na: the observed number of alleles; Ne: the effective number of alleles; H_o: observed heterozygosity; H_e: expected heterozygosity; I: Shannon’s information index.

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Figure 4. Development and evaluation of a core collection. (A) The minimized number of accessions for a core collection capturing the majority of alleles by random sampling and the Mstrat strategy. (B) Number of alleles observed in the total sample and those captured in the core collection, also presented separately for alleles occurring in different frequency. (C) NJ tree of the 115 core accessions. Values at the branches denote bootstrap values.

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Supporting Information

Table S1  Detail information of cucumber resources. (XLSX)

Table S2  Cucumber core collection. (XLSX)

Figure S1  Pictures of typical cucumber accessions from the three model based populations. CG1149, “Chinese Long”; CG4357, Southern China type; CG5786, Dutch greenhouse; CG6600, Pickling; CG9191, Xishuangbanna; CG8039, Indian (PDF)

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Author Contributions

Conceived and designed the experiments: SH. Performed the experiments: QS, IQ, DS, SZ, HS, JL, YS, XZ, JZ. Analyzed the data: JL, IQ, QS, ZZ, RT. Contributed reagents/materials/analysis tools: DS, QS, JQ, DS, SZ, GS, HL, JL, YS, XZ, JZ. Wrote the paper: JL, IQ, ZZ, SH.