Genetic diversity of cacao (*Theobroma cacao* L.) populations and agronomic traits for breeding

Dissertation

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Submitted by

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“El equilibrio empieza a dar su luz, cuando arriba en la montaña baila un guerrero noches enteras”.

-El equilibrio parte I, Jaguares
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PREFACE

This dissertation consists of a general introduction to cacao and the use of genetic resources for crop improvement, then three core research papers are presented, two published and the third submitted to an international peer-reviewed journal. It concludes with a general discussion that presents the main findings, conclusions, and research gaps related to conserving and using cacao genetic resources for crop breeding.

The first scientific paper, Geographic distribution, conservation, and genomic resources of cacao, *Theobroma cacao* L., analyses the geographic distribution of cacao and pinpoints new areas for germplasm collection. It discusses the conservation of cacao genetic resources, the genetic basis of agronomic traits, and genomic resources for cacao improvement.

The second paper, Geographic patterns of genetic variation among cacao (*Theobroma cacao* L.) populations based on chloroplast markers, focuses on the phylogeographic structure of natural cacao populations; chloroplast DNA (cpDNA) polymorphisms were used to show how chloroplast genetic diversity is geographically distributed in Amazonia, and how it is represented in common cacao genotypes of farms and breeding populations.

The third paper, Characterization of cacao (*Theobroma cacao* L.) accessions based on published SNP markers linked to agronomic traits reveals signals of selection, provides information on population structure of wild and cultivated cacao, identifies agronomic traits under selection, and uncovers valuable cacao genotypes not observed in cultivated and breeding populations.
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LIST OF ABBREVIATIONS

AMOVA Analysis of Molecular Variance
BPR Black Pod Rot
CATIE International Center for Tropical Agriculture
Cd Cadmium
CSE Caffeoyl-Shikimate Esterase
FAs Fatty Acids
FPR Frosty Pod Rot
GS Genomic Selection
GWAS Genome-Wide Association Studies
HMA Heavy Metal ATPase
ICGD International Cocoa Germplasm Database
ICGT International Cocoa Genebank Trinidad
LRR Leucine-Rich Repeat
NB Nucleotide-Binding
NRAMPS Natural Resistance-Associated Macrophage Proteins
PCoA Principal Coordinates Analysis
PR Pathogenesis-Related
QTL Quantitative Trait Loci
SC Self-Compatibility
SI Self-Incompatibility
SNP Single-Nucleotide Polymorphism
SSR Simple Sequence Repea;
cpSSR chloroplast SSR
WBD Witches’ Broom disease
GENERAL INTRODUCTION

1 THE CHOCOLATE TREE AND ITS GENETIC DIVERSITY

The cacao tree (*Theobroma cacao* L.), a diploid species (*2n = 2x = 20*) with a small genome (430–445 Mbp), belongs to the Malvaceae family and is native to Amazonia (Argout et al., 2011; Motamayor et al., 2013). Cacao is a shade-tolerant and relatively small tree. Its small, bisexual flowers are pollinated by midges, and the cauliflorous fruits have a hard and rugose surface, containing 20-40 seeds surrounded by sweet pulp (Wood & Lass, 1985). Cacao is cultivated in the tropics mainly by smallholder farmers under agroforestry systems to produce seeds, the unique source of chocolate (Coe & Coe, 2013; Daymond & Bekele, 2022; ICCO, 2021). A higher demand from the global chocolate industry drives increments in cacao cultivation globally, top producers include west Africa (76%), Latin America (17%) and Asia (6%) (FAOSTAT, 2020; ICCO, 2021). The global cacao production in 2020/21 was 5175 thousand tons of seeds (ICCO, 2021), major per capita consumers of chocolate include Europe and the US (Lindt & Sprüngli, 2018).

Western Amazonia is the hot spot of cacao genetic diversity. Seven out of eleven genetic clusters identified with nuclear microsatellite markers are located in the Ecuadorian-Peruvian Amazon (Motamayor et al., 2008; Zhang et al., 2012). This area also holds the highest variation of chloroplast haplotypes when compared to eastern Amazonia (Nieves-Orduña et al., 2023). Future collection trips likely will reveal new genetic clusters, due to high biological diversity of western Amazonia (Bass et al., 2010; Steege et al., 2003; Tuomisto et al., 1995). Unfortunately, due to persistent deforestation in Amazonia, the cacao primary gene pool is losing habitat, which likely is reducing the population size of natural populations (Smith & Schultes, 1990; Soares-Filho et al., 2006; Steege et al., 2015). Although there have been collection trips for wild material in Amazonia since the 1930s (Zhang & Motilal, 2016), areas
not yet explored for cacao *ex situ* collections with high probability of capturing new genetic diversity include river systems at the upper border between Ecuador-Peru (Nieves-Orduña et al., 2023). In addition, this area is important for collecting wild material that likely coevolved resistance to the damaging Frosty Pod Rot disease (caused by *Moniliophthora roreri*) (Díaz-Valderrama et al., 2022).

**2 DOMESTICATION AND BREEDING OF CACAO**

Domestication of cacao occurred ~3600 years ago in South America (Cornejo et al., 2018), with evidence of cacao use in Ecuador dating ~5600 years ago thanks to the recovery of ancient cacao DNA at the archaeological site Santa Ana – La Floridad of Ecuador (Zarrillo et al., 2018). The domestication of cacao, like other fruit tree species of Amazonia, likely targeted traits such as pulp flavor, which was consumed directly but also used for the preparation of fermented beverages (Clement et al., 2010; Venturieri, 2013). Traits such as reduced bitterness were selected by native Americans in ancient Criollo populations (Cornejo et al., 2018). In addition, cacao domestication led to the accumulation of deleterious mutations affecting fitness and yield in Criollo (Cornejo et al., 2018).

Criollo cacao was cultivated during colonial times, but modern cacao varieties are mainly hybrid varieties that preserve the traditional cacao flavor (Bartley, 2005; Cornejo et al., 2018). Populations genetic analysis based on 200 genomes revealed that cultivated cacao and breeding populations are mainly composed of hybrids with a high ancestry of Criollo and Amelonado clusters (Cornejo et al., 2018), confirming the underutilization of cacao genetic resources for developing breeding populations and cultivars (Zhang & Motilal, 2016). Since the beginning of the cacao trade in the seventeenth century to present day, the fine flavor attributed to Criollo ancestry has had a higher price at the international markets (Dand, 2011; ICCO, 2022).
The narrow genetic diversity of cultivated Criollo led to a series of historical reductions in cacao production likely due to the lack of disease resistance traits (Díaz-Valderrama et al., 2020; Zhang & Motilal, 2016). Examples include the cacao blast in 1727 in Trinidad, where mostly Criollo was cultivated, which forced the first introduction of wild cacao (called Forastero) to broaden the genetic base of cacao farms (Zhang et al., 2011). The first cacao breeding program took place in Trinidad and targeted yield and high seed weight while preserving the fine chocolate flavor (Toxopeus, 1969). Three old diseases destroyed up to 38% of global crop production and have been the focus of cacao breeding: Black Pod-BP (caused by Phytophthora spp.), Witches Broom Disease-WBD (caused by M. perniciosa), Frosty Pod Rot-FPR (caused by M. roreri) (Gutiérrez et al., 2016; Marelli et al., 2019). Traditional cross breeding has successfully preserved chocolate attributes while combining disease resistance and yield traits sourced from selected wild cacao (Boza et al., 2014; Goenaga et al., 2009; Phillips-Mora et al., 2013).

Modern genomic resources in cacao breeding include the analysis of two reference genomes Criollo B97-61 and Matina1-6, and characterization of key genes related to fat biosynthesis, fine flavor, and fruit color (Argout et al., 2011; Motamayor et al., 2013). The Infinium cacao 6K/15K SNP array is available for molecular breeding (Livingstone et al., 2015; Livingstone et al., 2017), the EMBRAPA multispecies chip, which includes 3.4K of these cacao SNPs, has made genotyping more affordable at around USD$20 per sample (Lopes et al., 2022). The cacao SNP array has been used in QTL mapping (Gutiérrez, Puig, et al., 2021; Royaert et al., 2016), GWAS studies (Romero-Navarro et al., 2017), genomic selection (McElroy et al., 2018) and estimation of genome-wide SNP homozygosity for developing inbred lines and exploiting heterosis (Lopes et al., 2022). In addition, cacao SNPs to facilitate the selection process are available for agronomic traits such as disease resistance to FPR, BP, and WBD (Gutiérrez, Puig, et al., 2021; Royaert et al., 2016), and yield (Bekele et al., 2022;
A set of 216 SNPs selected over the ten cacao chromosomes is available to classify and identify germplasm accessions (Gutiérrez, Martinez, et al., 2021).

Breeding in cacao still is a challenge but the primary gene pool holds novel alleles for present and future breeding activities (Zhang & Motilal, 2016). Traits of economic importance include disease resistance (especially to FPR), yield, reduced seed cadmium content, reduced tree height, sexual compatibility, and flavor (Bartley, 2005; Bekele & Phillips-Mora, 2019; Gutiérrez et al., 2016; Lanaud et al., 2017). Up to date the conservation of cacao genetic resources is organized in 40 ex situ collections (over 24k accessions are maintained), including two international collections (CacaoNet, 2012). In addition, challenges remain for the management of ex situ collections such as identification of duplicated and mislabeled accessions, genetic gaps, standardization of agronomic phenotyping, protocols for cryopreservation, and financial support to ensure long term conservation, where the chocolate industry can play an important role (CacaoNet, 2012; Nieves-Orduña et al., 2023).

Planting genetically improved varieties secures crop productivity and decreases pressure for new farmland, which reduces the risk of deforestation linked to cacao cultivation, and secures economic benefits for farmers and sustainability for the chocolate industry (Bekele & Phillips-Mora, 2019; Gutiérrez et al., 2016; Morrissey et al., 2019). Efficient conservation and use of underutilized cacao genetic resources are fundamental to address current and future breeding objectives.

This research aims to study the genetic diversity, conservation, and utilization of cacao genetic resources for crop improvement. Two types of genetic markers were used to study the genetic diversity of cacao populations (wild, cultivated, and breeding populations), cacao chloroplast microsatellite (simple sequence repeat - SSR) markers and published SNP (single nucleotide polymorphism) markers associated with economically important agronomic traits.
such as disease resistance and yield. A comprehensive review analyzing the conservation and utilization of cacao genomic resources for cacao breeding is also presented.

The specific objectives of this dissertation are:

- to identify the potential geographic distribution of wild cacao populations in Amazonia and show patterns of geographic genetic variation,
- to assess the conservation status and available cacao genomic resources for cultivar development,
- to review the genetic basis of agronomic traits such as disease resistance, quality traits and yield, and available molecular markers for cultivar development,
- to establish the geographic distribution of chloroplast (cpDNA) variation in different populations representing natural cacao stands, cultivated cacao, and breeding populations,
- to describe the population structure and analyze genetic clusters in wild and managed cacao, and identify underutilized genetic resources for cacao breeding,
- to identify agronomic traits under selection that differentiate wild and managed cacao to assist breeding of superior cacao genotypes.
3 REFERENCES


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Geographic distribution, conservation, and genomic resources of cacao 

*Theobroma cacao* L

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Geographic distribution, conservation, and genomic resources of cacao *Theobroma cacao* L

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**Core Ideas**

- The geographic distribution of genetic variation of wild cacao pinpoints new sampling areas for germplasm collections.

- The natural habitat of cacao is under threat due to current and projected deforestation in Amazonia.

- Germplasm collections can benefit from genotype identity and characterization of agronomic traits based on available SNP markers.

- The wide cacao genetic diversity can be tested against different fungal strains to identify new sources of disease resistance.

- Genome-wide SNP data in combination with standard phenotyping of agronomic traits can be exploited to identify relevant genotypes for breeding.
Abstract

The cacao tree, *Theobroma cacao* L., is cultivated in the tropics, mainly in agroforestry systems, to produce seeds, the valuable raw material for the chocolate industry. Thus, the conservation and use of cacao genetic resources in breeding programs to increase yield and improve quality and disease resistance are vital for the global cacao economy. We review three important topics of cacao genetics essential for sustainable production and crop improvement: (1) geographic distribution of wild cacao populations and geographic patterns of their genetic variation in Amazonia; (2) conservation, availability, and use of cacao genetic resources in cultivar development; (3) genetic basis of agronomic traits, available molecular genetic markers, and genomic resources and their application for cacao improvement. We also highlight critical research areas needed to achieve sustainable cacao cultivation.
1 INTRODUCTION

The cacao tree (*Theobroma cacao* L.) is a diploid (2n = 2x = 20) species with a comparatively small genome (430–445 Mbp) (Argout et al., 2011; Motamayor et al., 2013). It is a relatively small tree, up to 20 m high in natural forests, and shade-tolerant (Wood & Lass, 1985). Cacao is native to Amazonia and belongs to the Malvaceae family (Bayer & Kubitzki, 2003; Motamayor et al., 2002). The genus *Theobroma* includes 22 species native to tropical America, but none of these wild relatives has contributed to cacao improvement because of strong interspecific crossing barriers (Cuatrecasas, 1964; Zhang et al., 2011). The small and bisexual cacao flowers (developed in small inflorescences) are pollinated by midges of the Ceratopogonidae family (Young, 1994). The cauliflorous cacao fruits of globose to fusiform shape have a hard and rugose surface and exhibit a thick pericarp. On average, there are around 20–40 oblong seeds per fruit (pod) with white and sweet surrounding pulp (Cuatrecasas, 1964).

Cacao is cultivated in 61 tropical countries for the production of seeds, the raw material that shapes the world's chocolate industry (Coe & Coe, 2013; Dand, 2011; FAOSTAT, 2020). Per capita chocolate consumption is very high in both Europe (4–8 kg/year) and the USA (4 kg/year) (Lindt & Sprüngli, 2018). As a tree crop common to small farms and agroforestry systems, cacao seeds are commodities produced primarily in West Africa (76%), followed by Latin America (17%) and Asia (6%) (ICCO, 2021; Nair, 2021). The world's cacao production has been concentrated in West Africa since the 1980s, and top producing countries like Cote d'Ivoire and Ghana have experienced induced deforestation linked to cacao cultivation (Kroeger et al., 2017; Ruf & Schroth, 2004; Wessel & Quist-Wessel, 2015).

Globally, around 2–3 million hectares of tropical forest were lost due to cacao cultivation over a 20-year period (1988–2008) (Kroeger et al., 2017). Low yields, inadequate maintenance, pests and diseases, and increased international demand for chocolate are the main drivers of land use expansion and deforestation linked to cacao cultivation (Hoang & Kanemoto, 2021;
Kroeger et al., 2017; Morrissey et al., 2019; Wessel & Quist-Wessel, 2015). In addition, in West Africa, climate change scenarios predict a rise of 2°C during dry seasons by 2050, reducing suitable lands for cacao cultivation. These potential changes increase the risk of future deforestation in Liberia, Cameroon, and the Congo basin because cacao cultivation may be relocated to cooler areas (Schroth et al., 2016, 2017).

Cacao cultivation may play a dual role in Latin America. For example, it has contributed to an initial deforestation of the Atlantic Forest landscape but to its restoration later in the state of Bahia, Brazil (Bright & Sarin, 2004; Ruf & Schroth, 2004; Schroth et al., 2011). Similarly, both agroforestry reforestation and deforestation were linked to cacao farming in specific areas of Peru and Nicaragua (Orozco-Aguilar et al., 2021). Cacao farming was not associated with deforestation in Colombia but rather opposite—with agroforestry reforestation, which actually is an attractive alternative to illicit crops (Abbott et al., 2018; Barrera-Ramírez et al., 2019; Castro-Nunez et al., 2020; Hernández-Núñez et al., 2022).

As a shade-tolerant tree, more productive cacao can be planted together with valuable native trees creating an agroforestry that increases biodiversity by providing habitat to different plant and animal species (Bhagwat et al., 2008; Blaser et al., 2018; Jezeer et al., 2017; Rice & Greenberg, 2000; Schroth & Harvey, 2007). In addition, cacao agroforestry provides ecosystem services such as carbon capture and storage, water regulation, climate adaptation, and production of timber from shade trees (Blaser et al., 2018; Heming et al., 2022; Martin et al., 2020; Tscharntke et al., 2011). Compared to monoculture, cacao agroforestry with an intermediate level of shade (30%–50%) can equally be profitable (Blaser et al., 2018; Jezeer et al., 2017). Market recognition of biodiversity conservation values generated by cacao agroforestry can be achieved through certification schemes (Tscharntke et al., 2015).
Due to its importance in national and international markets, the genetic diversity of cacao trees has been studied to explore and improve the crop potential in agronomic traits, such as disease resistance, yield, climate resilience, and seed quality (Bekele & Phillips-Mora, 2019; Bennett, 2003; Daymond & Bekele, 2022; Dias, 2001; Gutiérrez et al., 2016; Hunter, 1990; Schnell et al., 2005). Here, we focus on the conservation and use of cacao genetic resources and the needs to be addressed in the face of current and future challenges.

2 GEOGRAPHIC DISTRIBUTION OF NATURAL POPULATIONS

The wide geographic distribution of cacao in the Amazon basin includes the tropical forests of Peru, Ecuador, Colombia, Bolivia, Brazil, Venezuela, and French Guyana (Figure 1). Cacao is observed from west to east Amazonia in non-flooded and seasonally flooded forests (e.g., high várzea forest, where the water level is less than 3 m in height and lasts less than 50 days/year) (Draper et al., 2019; Wittmann et al., 2006). A recent survey of Amazonia tree diversity observed cacao mostly in non-flooded forests (Steege et al., 2013, 2020). Groups of trees occur in neutral to alluvial soils also scattered and isolated in the understory of primary forests with a density of 1–10 individuals per ha (Allen, 1988; Cuatrecasas, 1964). More recent observations in Madre de Dios, Peru and Rondônia, Brazil showed higher densities, up to 85 and 142 trees/ha, respectively (Steege, 2021). These occurrences may have been influenced by patterns of human introductions in modified forests (Levis et al., 2017; Somarriba & Lachenaud, 2013).

The cacao tree occurs in areas with a mean annual temperature range of 20–30°C and precipitation ranging from 2000 to 6000 mm/year (Figure 2), at an altitude below 1000 m above sea level (masl) (Allen, 1988; Somarriba & Lachenaud, 2013). Recent observations in the South of Peru showed that cacao tolerates dry seasons and can grow at higher altitudes up to 1400 masl (Ceccarelli et al., 2021).
FIGURE 1. Potential geographic distribution of *Theobroma cacao* L., genetic clusters, and habitat loss observed in Amazonia. Light gray circles represent 4413 observations of cacao in wild habitats according to ecological studies and expedition reports for the 1938–2022 time period (Allen, 1988; Almeida, 1983; Almeida et al., 1987, 1995, 2015; Arevalo-Gardini et al., 2019; Barriga, 1982; Carletto, 1973; Ceccarelli et al., 2021; Chalmers, 1968, 1969, 1972, 1973; Chumacero de Schawe et al., 2013; Clement, 1986; Cocoa Research Unit, 1995; Coral, 1988; Desrosiers & Buchwald, 1961; Dias et al., 2003; Fouet et al., 2022; Garcia-Davila et al., 2020; González-Orozco et al., 2020; Lachenaud et al., 1997, 2016; Lachenaud & Sallée, 1993; Lanaud, 1986; Loor et al., 2015; Marita et al., 2001; Morales & Rodriguez, 1987; Motamayor et al., 2008; Pound, 1938, 1943; Sallée, 1987; Sanchez et al., 1988; Sereno et al., 2006; Soria, 1970; Steege, 2021; Turnbull & Hadley, 2021; Vásquez-García et al., 2022; Vello & Medeiros, 1966; Wittmann et al., 2006; Zhang et al., 2012). Observation records include data managed by the Amazon Tree Diversity Network (Steege, 2021) and the International Cocoa Germplasm Database (Turnbull & Hadley, 2021). Cacao genetic clusters are depicted by their respective symbols (circles, triangles, and squares of different colors) according to Motamayor et al. (2008), Zhang et al. (2012), and Gutiérrez, Martinez et al. (2021). The deep green layer shows forest cover (Karra et al., 2021), red polygons represent areas of increased deforestation (Pacheco et al., 2021), and the orange oval shows the border among Ecuador, Colombia, and Peru where cacao observation is low but expected diversity is high (Zarrillo et al., 2018). The map was generated using the ArcGIS software (www.esri.com).
2.1 Classification and primary genetic clusters

Morris (1882) (cited in Cuatrecasas, 1964) proposed a classification of cacao populations based on two morphological traits: color of fruits (red to yellow) and fruit surface (smooth or rough). Cacao was divided into two groups: Criollo (red fruits) and Forastero (consisting of eight types, yellow to red fruits and smooth and rough skins) (Morris, 1882). In general, Criollo is known for its smooth fruit skin and low seed productivity, and Forastero is characterized by a rough fruit surface and higher seed yield (Cuatrecasas, 1964). A third group, called Trinitario (Cheesman, 1944), a hybrid (Criollo × Forastero) originated in Trinidad and Tobago as a result of natural hybridization in cacao farms (Bartley, 2005; Yang et al., 2013). The classification as Criollo, Forastero, and Trinitario groups was traditionally used as a reference of geographical origin of the cacao populations: Forastero from the Amazon basin, Criollo from Central and
South America, and Trinitario from Trinidad, respectively (Bartley, 2005; Cuatrecasas, 1964). However, this initial classification does not reflect the rich genetic diversity of cacao populations observed in Amazonia, making the term Forastero obsolete (Almeida et al., 2015; Arevalo-Gardini et al., 2019; Cornejo et al., 2018; Loor et al., 2015; Motamayor et al., 2008; Motamayor et al., 2010; Zhang et al., 2009).

Motamayor et al. (2008) identified 10 clusters based on the analysis of 952 cacao accessions of widespread geographic origin conserved in *ex situ* collections with 96 simple sequence repeat (SSR) markers. The 10 clusters were named according to the geographic origin of accessions and their genetic clustering based on the SSR markers: Marañon, Curaray, Criollo, Iquitos, Nanay, Contamana, Amelonado, Purús, Nacional, and Guiana (Figures 1 and 2). Three clusters, Criollo, Nacional, and Amelonado, correspond to traditional cultivars. Morphologically, Criollo has plump seeds and white cotyledons; the remaining clusters, which can be grouped as Forastero, have flat seeds and purple cotyledons (Daymond & Bekele, 2022). Hence, this new classification based on cluster analysis of SSR genotypes provides more information about the available genetic resources for conservation and breeding programs (Laliberté et al., 2018; Motamayor et al., 2008; Zhang & Motilal, 2016). The SSR genotypes of 57 wild samples collected along the Beni River in Bolivia revealed an additional cluster, Cacao Nacional Boliviano (Zhang et al., 2012).

Recently, genotypes based on 219 single-nucleotide polymorphisms (SNPs) validated the SSR-based clusters observed in Amazonia and provided higher resolution to study populations and subpopulations within the primary gene pool of cacao in Amazonia (Gutiérrez, Martinez, et al., 2021). For example, accessions from the Napo River in Ecuador assigned to the Purús cluster with a low support in the SSR study were firmly assigned to this cluster based on the SNP genotypes. Moreover, samples geographically restricted to the Purús River in Brazil formed a new cluster—population Purús II—separate from the original Purús cluster. In
addition, the Iquitos cluster is likely restricted to the Peruvian Amazon based on the SNP genotypes, excluding samples collected in Central Amazonia (Gutiérrez, Martinez, et al., 2021). Thus, the 219 SNP panel can be used to classify cacao germplasm more accurately. Overall, SNP genotypes from accessions conserved in national cacao collections may reveal unique ancestry based on geographic origin.

Collecting and analyzing different wild populations in Amazonia with total estimated number of 180 million cacao trees (Steege et al., 2020) might reveal novel cacao genetic diversity and clusters (Bartley, 2005; Laliberté et al., 2018; Zhang et al., 2009; Zhang & Motilal, 2016). For example, wild cacao was observed in northern Amazonia (Figure 1) (Colombia, Venezuela, and Brazil), but it was not studied yet, and no genetic clusters were reported for these areas. Similarly, important river systems in northwestern Amazonia like the upper Napo, Tigre, Pastaza, and Morona in Peruvian Amazon could host new genetic diversity (area depicted by the orange oval in Figure 1), especially considering the frequent observation of wild cacao and occurrence of five genetic clusters in the upper Amazon (Figure 1).

2.2 Geographic origin of clusters
The geographic origin of cacao genetic diversity in Amazonia is attributed to paleoclimate and paleoarches (Motamayor et al., 2008). The paleoarches were ridges no longer visible in the landscape that shaped the genetic structure of cacao by acting as gene flow barriers between cacao clusters (Motamayor et al., 2008). The paleoclimate refers to past dry and cool conditions during the Quaternary period in Amazonia (van der Hammen & Hooghiemstra, 2000). According to the refugia theory, forests were reduced to islands or refugial areas, where temperature, humidity, and edaphic conditions protected tree growth and diversity and promoted genetic differentiation between populations due to geographic isolation (Gentry, 1982; Haffer & Prance, 2001; Prance, 1973). In addition, present-day adaptation of tree species to gradients of soil fertility in Amazonia also explains the occurrence of tree populations in
specific habitats with interrupted gene flow (Gentry, 1989). In cacao, the geographic
distribution of genetic diversity is not only attributed to a positive response to fertile soils but
also to the geographic isolation of populations growing along different Amazonian rivers
system, where gene flow among populations is limited, favoring genetic differentiation

Based on the rich diversity of crop species observed in Amazonia and putative forest
refugia, a center of crop genetic diversity for fruit tree species, including cacao, was proposed
in western Amazonia (Clement, 1989, 1999). Lachenaud (1997) suggested the existence of
forest refugia during the Pleistocene epoch to explain the actual heterogeneity and
discontinuous distribution of wild cacao in Amazonia. Likewise, five forest refugia for cacao
in Amazonia were proposed by modeling potential past geographic distributions to explain the
putative origin of cacao clusters (Thomas et al., 2012). The rich chloroplast diversity observed
in wild cacao compared to cultivated cacao supports the forest refugia theory in northwestern
Amazonia (Nieves-Orduña et al., 2021). Furthermore, the distribution of cacao genetic
diversity in Amazonia may have also been affected by ancient and modern human intervention
(Levis et al., 2017; Peters, 2000; Somarriba & Lachenaud, 2013; Thomas et al., 2012), which
may explain the convergence of five genetic clusters (Iquitos, Nanay, Marañon, Contamana,
and Purús) in the Peruvian Amazon, and the patterns of diversity decline from west to east in
Amazonia (Figure 1; Cornejo et al., 2018; Motamayor et al., 2008; Nieves-Orduña et al., 2021).

3 EX SITU AND IN SITU CONSERVATION OF CACAO

3.1 Cacao expeditions and germplasm collections

After the collapse of cacao plantations in Trinidad around 1930 as a result of witches’ broom
disease (WBD) caused by Moniliophilthora perniciosa, there was a need to look for new sources
of genetic resistance in natural populations of Amazonia (Bartley, 2005). The narrow genetic
base of cultivated cacao also became evident later on in Bahia, Brazil, where in the 1990s the production decreased dramatically due to the vulnerability to WBD (Bennett, 2003). Thus, cacao germplasm collections in Amazonia have been motivated by the search for disease resistance and yield traits (Bartley, 2005; Zhang et al., 2011). The search for disease resistant genotypes focused extensively on the northern Peruvian Amazon by the collector F.J. Pound during the 1930–1940s (Bartley, 2005; Zhang et al., 2011). More recently, systematic field trips aiming to collect genotypes expressing disease resistance and superior yield traits searched wild cacao trees in the Colombian and Ecuadorian Amazon (Allen, 1988). Detailed descriptions of historical cacao expeditions collecting material for ex situ collections are presented in Zhang et al. (2011), Zhang and Motilal (2016), and Rodriguez-Medina et al. (2019).

New cacao expeditions were motivated to widen the genetic base for cacao breeding and filling gaps in ex situ collections (Bekele & Phillips-Mora, 2019; Daymond & Bekele, 2022; Laliberté et al., 2018). Collection trips aiming to capture new genetic diversity for disease resistance and quality traits were done in Peru (Arevalo-Gardini et al., 2019; Vásquez-García et al., 2022; Zhang & Motilal, 2016), Ecuador (Fouet et al., 2022; Loor et al., 2015), and French Guiana (Lachenaud et al., 2018). In addition, not only disease resistance but also quality traits have been subject of specific cacao collections. In northern Peru, collections of wild cacao were conducted to investigate populations of the fine flavored cacao variety “Piura Porcelana” (Arevalo-Gardini et al., 2019). Specific collections aiming to rescue and conserve wild ancestors of the fine flavored Nacional group were conducted in the Ecuadorian Amazon (Fouet et al., 2022; Loor et al., 2015). In addition, expeditions in the Colombian Amazon along the Caquetá and Caguan rivers identified wild cacao populations and potential areas for germplasm collection (González-Orozco et al., 2020; Rodriguez-Medina et al., 2019).
New germplasm collections could focus on areas where expected genetic diversity is high (Figure 1), such as northwestern Amazon (Motamayor et al., 2008; Nieves-Orduña et al., 2021; Thomas et al., 2012; Zarrillo et al., 2018). However, germplasm collections have to be done on time before deforestation and climate change erodes the natural habitat of cacao in Amazonia (Gomes et al., 2019; Soares-Filho et al., 2006).

3.2 *Ex situ* conservation

Cacao needs to be conserved as a living plant. Due to the recalcitrant nature of cacao seeds, they can be conserved only up to 10 weeks and do not tolerate desiccation below 27% and temperatures lower than 17°C (Hor et al., 1984). Hence, cacao needs to be conserved as living collections of trees, thus each tree represents a unique genotype in germplasm collections (CacaoNet, 2012). The *ex situ* conservation of cacao genetic diversity is currently arranged mainly via 47 national collections and two international collections under the public domain regulated by the International Treaty on Plant Genetic Resources for Food and Agriculture (Motilal, 2018; Thormann et al., 2015). International collections are maintained in Costa Rica at CATIE (International Center for Tropical Agriculture) and Trinidad and Tobago at ICGT (International Cocoa Genebank Trinidad, West Indies University). Accessions maintained in CATIE and ICGT collections represent wild, breeding populations (mostly as admixture trees), and genetic clusters. Among the genetic clusters conserved, Marañon, Nanay, Iquitos, and Amelonado are the most frequent, Criollo is the least represented, and Cacao Nacional Boliviano is absent.

A core germplasm collection aims to capture the total crop genetic diversity by including an optimal set of accessions for trait evaluation and providing access for breeders (Maxted et al., 2020). Two hundred sixty-one germplasm accessions were proposed for a cacao core collection, including the 10 genetic clusters identified by Motamayor et al. (2008) plus 11 Trinitario accessions (CacaoNet, 2012; Motilal, 2018). The criteria for germplasm selection...
did not include agronomical or morphological trait information, but individual SSR and geographic genetic variation data observed in different ex situ collections (CacaoNet, 2012). In the case of Guianan cacao, through the analysis of 1953 SNPs, 41 out of 181 cacao clones were selected as a core collection, but 21 clones captured 99% of all alleles in a core collection (Lachenaud et al., 2018).

Cacao is conserved also in national germplasm ex situ collections mainly in countries with natural cacao populations such as Brazil (4773 accessions), Ecuador (1939), Peru (593), Colombia (411), Venezuela (419), and French Guiana (551) (Turnbull & Hadley, 2021). Among them, Brazil and Ecuador are top cacao producing countries in Latin America (ICCO, 2021). Although cacao is native to Bolivia, there are no records of ex situ conservation in this country (Turnbull & Hadley, 2021). The west African countries, which produce 76% of global cacao (ICCO, 2021), also hold cacao accessions, for example, Cote d'Ivoire (411), Ghana (1278), and Cameroon (230) (Turnbull & Hadley, 2021). In total, around 18,634 cacao germplasm accessions are conserved in national ex situ collections (Turnbull & Hadley, 2021). When considering specific national regulations, breeders may face challenges to access and exchange cacao accessions due the exclusion of cacao from Annex I of the International Treaty on Plant Genetic Resources for Food and Agriculture. As a response, the Global Strategy for cacao genetic resources aims to improve conservation and access of cacao germplasm through the creation of a network of international partners, including countries that maintain ex situ collections, national authorities, and the private sector like chocolate companies (CacaoNet, 2012; Laliberté et al., 2018).

Agronomic evaluation carried out in ex situ collections has been fundamental for selecting commercial cacao clones and their incorporation in breeding programs (Bekele & Phillips-Mora, 2019). For example, in Trinidad and Tobago, among 1900 diverse cacao accessions conserved and evaluated at the ICGT, 57 Trinitario clones were recommended for cacao
plantations and as parents in breeding programs for their favorable yield potential, bean quality, and chocolate flavor (Bekele et al., 2020). Trinitario accessions have an impact on cacao producing countries worldwide (Bartley, 2005). Distributed in Latin America, Asia, and Africa, around 36 common Trinitario accessions (e.g., ICS and UF clones) have been used as planting material and breeding accessions (Bekele et al., 2020; Bekele & Phillips-Mora, 2019; Zhang et al., 2011). In French Guiana, CIRAD recommended three clones (known as GU clones) because of their disease resistance against WBD and black pod rot (BPR), yield performance, and intense chocolate flavor (Lachenaud et al., 2007). The agronomic evaluation of GU clones lasted 10 years as an initial population of 1600 wild cacao trees was collected in Guianan forests (Lachenaud et al., 2007).

Similarly, after 15 years of field trials, six clones with high yield and disease resistance to monilia or frosty pod rot (FPR) disease (caused by Moniliophthora roreri) and BPR disease were selected in the breeding program at CATIE in Costa Rica (Phillips-Mora et al., 2012). These selected clones (e.g., CATIE R6) have been widely distributed to cacao farmers in Central America and Brazil (Phillips-Mora et al., 2017). In addition, nine superior hybrids (TARS series) were selected after 4 years of evaluating yield and disease resistance in the germplasm collection maintained by the USDA in Puerto Rico (Goenaga et al., 2009). These 9 cultivars originated from a breeding population of 1310 trees representing 5 full-sib families (UF-668 × Pound-7, IMC-67 × UF-613, EET-400 × SCA-12, SCA-6 × EET-62, and IMC-67 × SCA-12) (Goenaga et al., 2009). Each clonal parent was previously selected in CATIE for its high yield and disease resistance traits (Goenaga et al., 2009). Further selections in Puerto Rico yielded three new cultivars (Gasic et al., 2016). Recent agronomic evaluation of native cacao from Peru identified 24 out 103 accessions with potential for the chocolate industry (Vásquez-García et al., 2022).
The cacao germplasm data are stored, and their access is facilitated by the International Cocoa Germplasm Database (ICGD, http://www.icgd.rdg.ac.uk) (Turnbull & Hadley, 2021). The ICGD provides valuable primary information on germplasm availability and agronomic traits for the research community, industry, and cacao growers (Daymond & Bekele, 2022). By collecting, updating, and managing germplasm information, the ICGD supports the selection of cacao accessions for conservation and breeding purposes. For example, breeders interested in cacao clones with confirmed resistance to FPR disease can check agronomic data of 18 cacao clones, including geographic origin, pedigree (when available), and the studies that support the selection of these disease resistant clones (Turnbull & Hadley, 2021).

3.2.1 Cryopreservation

Cryopreservation is an alternative to living collections for tree crops with recalcitrant seeds. In vitro material can be conserved at −196°C without losing viability for many plants (O'Brien et al., 2021). Reference germplasm collections can be preserved safely and cost-effectively in liquid nitrogen tanks when standard cryopreservation protocols are available (Jenderek & Reed, 2017). Cryopreservation of cacao somatic embryos was developed as a long-term alternative for genetic conservation (Adu-Gyamfi & Wetten, 2012, 2020; Fang et al., 2004). Cacao cryopreservation relies on floral-derived secondary somatic embryos because there are no methods for micropropagation of cacao shoot tips (Adu-Gyamfi & Wetten, 2020). Cryopreservation has been tested in six cacao genotypes, where the cultivar CCN51 exhibited a lower survival rate of 36% (Adu-Gyamfi & Wetten, 2012). However, to date, there are no cacao cryo-banks as a backup of ex situ collections. It seems that more research is needed to optimize and implement cacao cryopreservation protocols (vitrification) for specific genotypes (like cultivars and cacao core germplasm accessions). In addition, cacao embryogenesis, which tends to be genotype dependent, also needs to be validated in accessions of interest for cryopreservation (Wickramasuriya & Dunwell, 2018). Other research topics in developing
cryopreservation protocols should include studies of epigenetic and genetic changes and field performance of plants regenerated after cryopreservation (Wang et al., 2021). In cacao, phenotypic variability was observed in the cryopreserved accession AMAZ 15 and could be due to DNA methylation changes (Adu-Gyamfi et al., 2016).

3.3 In situ conservation

The natural habitats that host not only the primary genetic pool of cacao but also other 6000 tree species are under threat because of the continued destruction of Amazonian forests (Cardoso et al., 2017; Gomes et al., 2019; Rojas et al., 2021; Steege et al., 2015; Vancutsem et al., 2021). Unfortunately, it is projected that more than 40% of Amazonia will be lost by 2050 because of agricultural expansion and climate change (Gomes et al., 2019; Soares-Filho et al., 2006). The in situ conservation of cacao secured by national protected areas in Amazonia is experiencing different levels of threats due to illegal mining and logging, fires, and agricultural expansion (Armenteras et al., 2021; RAISG, 2020; Soares-Filho et al., 2006). Recently, five areas were proposed for in situ conservation and germplasm collection of wild cacao in the Colombian Amazon, and areas for cacao wild relatives (Theobroma spp. and Herrania spp.) were also identified in this study (González-Orozco et al., 2020).

Conservation strategies for widespread tropical tree species, such as cedar or cedro amargo (Cedrela odorata), have been proposed based on the distribution of cpDNA variants (haplotypes) in its natural habitat (Cavers et al., 2004). In the case of cacao, a center of cpDNA diversity was observed in the northwestern Amazon, including the occurrence of seven geographically restricted chloroplast haplotypes in the Peruvian Amazon (Nieves-Orduña et al., 2021). Because cacao is a non-pioneer species, in situ conservation needs to consider ecological conditions provided by natural forests to favor seed germination and adaptive changes (Akinagbe et al., 2019; Finkeldey & Hattemer, 2007). However, research projects investigating the in situ conservation of cacao are scarce (Daymond & Bekele, 2022). The study
of cacao conservation units depends on the genetic variation present in Amazonia; therefore, an initial step should include population genetic analysis of wild populations conserved in national *ex situ* collections. In addition, novel genetic diversity in natural populations sampled in new collections trips should be studied and used in breeding programs. It is also important to study population structure, mating system, gene flow, and ancestry of cacao populations using selectively neutral cacao markers, such as SSRs and SNPs (Gutiérrez, Martinez, et al., 2021; Mahabir et al., 2020; Motamayor et al., 2008; Wang et al., 2020; Wever et al., 2019).

**4 GENETIC BASIS OF AGRONOMIC TRAITS**

Cacao breeding for chocolate production is still a challenge that requires the integration of data on the best agronomic traits obtained from different populations with modern genomic and breeding analytical and experimental tools. Figure 3 demonstrates how breeding process can profit from combining standard phenotyping with studies such as genome-wide association studies (GWAS) for determining the genetic basis of economic traits (right part of the figure). Pre-breeding stages include also accurate genotype identification of germplasm collections and selection of agronomic traits based on available SNP markers (left part of the figure). Marker-assisted selection, genomic selection (GS) to predict phenotype performance of new genotypes, and field trials can be used together to validate new genotypes before commercial planting (center part of the figure). Overall, the genetic basis of agronomic traits relies on the conservation and use of cacao genetic diversity. For example, the study of complex traits among different cacao populations (Figure 1) aided with GWAS can identify novel and useful alleles for breeding disease resistant and low cadmium (Cd) genotypes (Figure 3). Breeding programs can obtain pathogen-free cacao plants (typically budwoods) from the International Cocoa Quarantine Center in the United Kingdom (Daymond, 2018). However, access to more diverse populations is often limited due to phytosanitary control and genetic resources’ protective regulations in countries with natural cacao populations and *ex situ* collections.
(Laliberté et al., 2018). Overall, increasing the number of geographically distant individuals (e.g., up to 1000 in GWAS by da Silva et al., 2016) representing a wide range of phenotypic variation genotyped with a high-density SNP array (e.g., up to 100 K) can reveal novel marker-trait associations.

**FIGURE 3.** Use of genetic markers and modern genomic tools in combination with studies of agronomic traits for cacao breeding. Cacao fruit photos courtesy of USDA ARS.

In addition, standardization of phenotyping of economically important traits is essential in GWAS experiments. For example, the evaluation of FPR disease resistance over 4 years proved that quantitative internal disease measures, such as percentage of necrosis in fruits, were more accurate than external qualitative disease evaluation (Gutierrez et al., 2021). Thus, measurements of internal disease severity are better suited to select FPR resistance genotypes (Gutierrez et al., 2021). In addition, reduced yield variability from the same clone can be achieved by measuring seed weight from fruits manually pollinated or using the number of seeds in the fruits as a covariable (Doaré et al., 2020). Either option will normalize seed weight phenotyping, providing data comparability among breeding programs (Doaré et al., 2020).
Validation of candidate SNPs identified by GWAS under different plantation environments is also needed, although it may represent a challenge in cacao breeding.

4.1 Genetic basis of disease resistance

Diseases are common in cacao and destroy up to 38% of the annual global yield, badly affecting farmers, and the chocolate industry (Bowers et al., 2001; Marelli et al., 2019; Ploetz, 2016); with world market revenue estimated at 980 billion in 2021 (Statista, 2022). The top main cacao diseases affecting cacao cultivation are BPR caused by seven species of Phytophthora sp., WBD caused by *M. perniciosa*, cacao swollen shoot virus caused by Badnavirus, and FPR caused by *M. roreri* (Bowers et al., 2001; Gutiérrez et al., 2016; Marelli et al., 2019). The world chocolate economy will be at risk, if FPR and WBD, which currently are geographically restricted to tropical America, would be introduced to West Africa, where more than 70% of the global production relies on disease susceptible Amelonado varieties (Díaz-Valderrama et al., 2020; Marelli et al., 2019; Ploetz, 2016).

4.1.1 Black pod rot disease (BPR)

Globally, BPR is the most economically harmful cacao disease, especially in Africa, where yield losses can be up to 90%, if not controlled (Morales-Cruz et al., 2020). It is caused by two significant pathogens: *Phytophthora palmivora* (native to Southeast Asia) and more aggressive *Phytophthora megakarya* (native to Central Africa and confined to Central and West Africa) (Morales-Cruz et al., 2020). Some cacao natural populations and cultivars show resistance to BPR, as well as disease resistance candidate genes are also identified (Table 1). For example, the Guiana group showed natural resistance to *P. megakarya*: Disease resistance was evaluated using artificial inoculation with *P. megakarya* in detached cacao leaves (Paulin et al., 2008), and the resistance of Guiana progenies to BPR was confirmed in field evaluations in Ghana (Ofori et al., 2020). Field evaluation of 148 individuals with admixed ancestry showed that trees with higher Nacional ancestry were more susceptible to BPR, whereas those with
## Table 1. Disease resistance candidate genes for prospective genetic improvement of cacao *Theobroma cacao* L.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Chromosome</th>
<th>Molecular description</th>
<th>Reference</th>
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Marañón ancestry showed resistance to BPR (Romero-Navarro et al., 2017). Furthermore, inoculation of 60 different cacao genotypes identified 22 accessions with resistance to BPR, of
which 4 exhibited high disease resistance in some accessions in Nanay (2 accessions), Iquitos (1 accession), and Marañon (1 accession) genetic groups (Fister et al., 2020).

Using published SSR markers and field response to BPR of traditional cacao varieties (Comum, Pará, and Maranhão) cultivated in Bahia, Brazil, 160 candidate genes were identified as associated with resistance to BPR (Mucherino Muñoz et al., 2021). The candidate genes include RLK (receptor-like kinase domain) and RLK + GNK2 (ginkbilobin-2), coiled-coils, and nucleotide-binding sites. From the breeder's perspective, four SSR markers were associated with BPR resistance, explaining 1.93%–7.43% of the phenotypic variation (Mucherino Muñoz et al., 2021). Gene expression analysis of leaves inoculated with *P. palmivora* revealed higher expression levels of disease resistance genes in the BPR resistant genotype Pound 7 compared to common cacao cultivars ICS 1 and CCN 51 (Baruah et al., 2021). The differentially expressed genes included genes of the KEGG pathway (phenylpropanoid biosynthesis), CAZy families, ubiquitin related proteins, LRR (leucine-rich repeat), and NB-ARC (nucleotide-binding adaptor) (Baruah et al., 2021). Furthermore, gene expression analysis revealed 67 pathogenesis-related (PR) genes when cacao seedlings were infected with *P. palmivora* (Fister et al., 2016). Although the cacao PR gene family did not stop the infection (after 72 h of inoculation, the pathogen was re-isolated to prove successful infection), it demonstrates that PR genes in cacao are involved in leaf tissue defense against BPR (Fister et al., 2016). In addition, a pioneer study in cacao using the CRISPR/Cas9 strategy demonstrated that the inactivation of non-expressor of PR 3 (TcNPR3) gene suppressed cacao disease response, and deletions in the TcNPR3 gene in detached leaves showed more resistance to BPR (Fister et al., 2018). Another gene involved in cacao resistance to BPR includes the caffeoyl-shikimate esterase (CSE) gene, which synthesizes caffeic, an inhibitor of *P. palmivora* (Winters, 2022). Consistent upregulation of TcCSE gene expression in response to *P. palmivora* infection was observed across BPR resistant genotypes of Guiana, Iquitos, Marañon, and Nanay groups.
(Winters, 2022). Other BPR disease resistance candidate genes identified across five species of *Theobroma* that also showed upregulated gene expression in response to *P. palmivora* infection were TcWRKY29 (WRKY transcription factor 29) and TcBBE8 (berberine bridge enzyme 8) (Winters, 2022).

These findings provide evidence of natural resistance to BPR among wild cacao populations. Incorporation of disease resistance while including additional traits, such as higher yield in single genotypes, still remains a challenge for breeding programs, but progress has been made in West Africa (Ofori et al., 2020) and Tropical America (Gutiérrez et al., 2016; Gutiérrez, Puig, et al., 2021; Phillips-Mora et al., 2012).

4.1.2 Witches broom disease (WBD)

WBD caused by *M. perniciosa* is one of the most critical cacao diseases, particularly in Brazil, but, fortunately, wild cacao populations exhibit some natural resistance. Restricted to tropical America, and native to Amazonia, the range of hosts of *M. perniciosa* includes *Theobroma* spp. and trees from the Solanaceae family (Lisboa et al., 2020). Cacao yield losses can be very high when WBD occurs together with FPR (Marelli et al., 2019; Ploetz, 2007). Resistance to WBD was observed in wild cacao populations, such as C SUL and SCA accessions, both collected in the upper Amazon and used by breeding programs in Brazil (Bennett, 2003; Marita et al., 2001; Paim et al., 2006; Pimenta Neto et al., 2018). However, because *M. perniciosa* in Brazil could overcome the disease resistance of SCA 6, the primary source of disease resistance, new resistant cultivars are needed to sustain the cacao production (Pereira et al., 2021). Thus, field evaluations of WBD in Brazil using 11 cacao genotypes with wide geographic origin identified the wild cacao accessions C SUL-3 (collected in Western Amazon, Brazil) and GU-171 (collected in French Guyana) as novel sources for the WBD resistance (Pereira et al., 2021).
The genetic basis of cacao resistance to WBD is polygenic (Table 1). The analysis of progeny in the mapping population from the TSH 1188 × CCN 51 cross identified seven quantitative trait loci (QTL) with small effects on the WBD resistance (Royaert et al., 2016). The major QTL on chromosome 9 originated from the SCA 6 accession. Candidate genes were validated through expression data, and SNP markers with highest association with disease resistance were identified in seven candidate genes (Royaert et al., 2016). Recently, WBD resistance was also studied in the F2 population derived from the SCA 6 × ICS1 cross, and 23 QTL with minor effects were identified (Chia Wong et al., 2022). Three identified candidate genes encoding glutathione peroxidases, threonine–serine receptors, and endochitinases, respectively, were associated with WBD resistance (Chia Wong et al., 2022). Cultivated to produce pulp and seeds, the cupuassu tree (*Theobroma grandiflorum*) is also infected by *M. perniciosa*. The WBD disease resistance gene TgPR3 identified in cupuassu is orthologous to the cacao gene Tc06v2_p000370 (Mournet et al., 2020). The latter confers disease resistance against BPD, thus, validating these two genes as part of defense mechanisms against pathogens in two *Theobroma* species (Mournet et al., 2020).

Breeding for disease resistance to WBD should consider the geographic and genotypic variation of *M. perniciosa*, as resistant clones identified in Brazil and Trinidad were susceptible in Ecuador (Meinhardt et al., 2008). Indeed, *M. perniciosa* has a wide natural geographic range in Brazil (Lisboa et al., 2020), and its population structure revealed dispersal patterns from west Amazonia (isolates collected in wild forests) to the eastern cacao plantations of Bahia (Artero et al., 2017). Furthermore, three *M. perniciosa* genetic clusters without clear host specificity were observed in five cacao production areas of Colombia (Jaimes et al., 2022). Thus, the population structure of *M. perniciosa* was shaped by geographic origin and environmental differences rather than by host specificity (Jaimes et al., 2022). Overall, geographically different *M. perniciosa* isolates obtained from wild and cultivated cacao aided
by additional genetic markers (e.g., SSRs and SNPs) will inform about the population structure and distribution of *M. perniciosa* in cacao production regions of tropical America. This insight will be of value to national breeding programs studying host specificity and resistance of cacao populations against different WBD strains.

4.1.3 Frosty pod rot (FPR) disease

FPR caused by *M. roreri* is the most severe cacao disease, which is restricted to tropical America, excluding Brazil. Yield losses due to FPR can be up to 80%, forcing land use change or “boom-bust” cycles (Zhang & Motilal, 2016). It is a specialized pathogen that attacks only *Theobroma* spp. and *Herrania* sp. trees. Until recently *M. roreri* was considered native to the upper Magdalena River in Colombia, outside the natural geographic distribution of cacao (Ali et al., 2015; Phillips-Mora et al., 2007). However, recent population analysis of *M. roreri* based on SSR and SNP markers identified six genetic clusters in areas of intense cultivation in tropical America: Four clusters represented Coastal Ecuador and three—Magdalena Valley in Colombia, and one cluster included both countries (Díaz-Valderrama et al., 2022). Another study based on SSR markers identified high genetic variation in *M. roreri* from the Ecuadorian Amazon where three clusters were identified (Espinoza-Lozano et al., 2022). A comparison of *M. roreri* collected from plantations in coastal Ecuador and a cluster representing Amazonia suggests the introduction of *M. roreri* from the Ecuadorian Amazon to Coastal Ecuador (Espinoza-Lozano et al., 2022).

The putative geographic origin of *M. roreri* is the Upper Amazon, an area shared by Colombia, Ecuador, and Peru (Díaz-Valderrama et al., 2022). Likely, untapped cacao genetic diversity evolved in the presence of different variants of *M. roreri* in the upper Amazon (Figure 1). Indeed, a recent study shows that the accessions EBC 6 and EBC 9, collected in the Colombian Amazon (Allen, 1988), are resistant to FPR in Colombia (Osorio-Guarín et al.,
Further cacao collections should target the upper Amazon (e.g., the border between Ecuador and Peru) to collect cacao genotypes with new genes conferring resistance to FPR.

Cacao natural resistance to FPR has been identified in germplasm collections and plantations, but it is not common (Bailey et al., 2018; Phillips-Mora & Wilkinson, 2007). FPR artificial inoculation of fruits allowed to test the tolerance reaction of diverse cacao accessions against seven FPR isolates (Phillips-Mora et al., 2005). Using this approach on 746 cacao accessions with geographically diverse origin maintained at CATIE revealed limited FPR resistance: 2% of the accessions were classified as resistant and 8% as moderately resistant (Phillips-Mora et al., 2009). However, follow-up studies at CATIE identified 50 diverse clones with moderate resistance to FPR and one resistant clone, the artificial hybrid CATIE R6 (UF-273 T1 × PA-169) (Phillips-Mora et al., 2017). In addition, field evaluations of UF 273 × Pa 169 progenies in Mexico showed a 41.27% reduction in FPR incidence compared to the average phenotypic value of progeny trials that included 22 full-sib families (Bonilla et al., 2021).

Cacao resistance to FPR should consider the pathogen variability observed in different locations. For example, a recent study of FRP diversity in tropical America (including 228 isolates obtained from cacao farms in 6 countries) used SSR and SNP markers to detect 6 hypothetical ancestral populations of _M. roreri_ (Díaz-Valderrama et al., 2022). Two hotspots of _M. roreri_ genetic diversity were observed in Colombian and Ecuadorian cacao plantations (Díaz-Valderrama et al., 2022). Thus, cacao clones should be tested for disease resistance in different countries while considering pathogen variability (Gutiérrez et al., 2016). In addition, preventive breeding in cultivated areas where FPR is not yet present, such as west Africa, is necessary (Gutiérrez et al., 2016).
Ecuador and Colombia are ideal countries for breeding resistance to FPR, considering the introduction of different strains of *M. roreri* in areas of intense cacao cultivation (Díaz-Valderrama et al., 2022; Espinoza-Lozano et al., 2022; Jaimes et al., 2016). For example, artificial inoculation of five cultivated clones with seven isolates of *M. roreri* in Colombia identified the hybrid ICS 95 (Amelonado × Criollo) as a resistant clone (Phillips-Mora et al., 2005). This clone and its derivatives (e.g., CCN 51) are a common source of resistance to FPR in Latin America (Bailey et al., 2018). In addition, agronomic evaluations of diverse clones in cultivated areas of Colombia identified several cacao clones with resistance to FPR (Cubillos, 2017; Jaimes et al., 2019; Osorio-Guarín et al., 2020). Evaluation of the breeding population representing CCN 51 × EET Nacional-type clones in coastal Ecuador identified four high yield clones resistant to FPR, which also exhibited resistance to WB and BPR (Jaimez et al., 2020). Nacional admixture genotypes showed resistance to FPR also in Ecuador (McElroy et al., 2018). Unless disease resistant clones are delivered to farmers, managing that FPR relies on extra management practices, such as removing infected pods from plantations (Cubillos, 2017; Phillips-Mora et al., 2012).

4.1.4 Genomic resources for breeding disease resistance

GWAS studies and QTL analysis in cacao have identified different loci associated with resistance to FPR, BPR, and WBD (Gutiérrez, Puig, et al., 2021; McElroy et al., 2018; Osorio-Guarín et al., 2020; Romero-Navarro et al., 2017). In Costa Rica at CATIE, six genes located on chromosomes 4, 5, 6, and 10 were associated with FPR resistance (Romero-Navarro et al., 2017). A higher proportion of Nacional ancestry within the 148 accessions analyzed was associated with resistance to FPR but susceptibility to BPR (Romero-Navarro et al., 2017). Follow-up analysis at CATIE using artificial inoculation in the F1 population from the Pound 7 × UF 273 cross identified six QTL associated with resistance to FPR (Gutiérrez, Puig, et al., 2021). Six SNPs located in chromosomes 2, 4, 7, 8, 9, and 10 accounted for 52.6% of
phenotypic variation (Gutiérrez, Puig, et al., 2021). The same study identified four QTL associated with resistance to BPR; four SNPs located on chromosomes 2, 4, 8, and 10 accounted for 50.6% of the phenotypic variation (Gutiérrez, Puig, et al., 2021).

In Ecuador, the natural occurrence of FPR and WB was observed among Nacional admixture genotypes, seven SNPs associated with FPR resistance located on chromosomes 1, 3, 5, 9, and 10, and nine SNPs associated with WBD resistance located on chromosomes 1, 2, 6, 7, and 9 were identified as markers for GS, which demonstrated a higher predictive accuracy of disease resistant plants when compared to single-marker selection (McElroy et al., 2018). For the cacao germplasm collection hosted in Colombia, two SNP markers were associated with FPR resistance on chromosomes 1 and 2, respectively, and four with WBD resistance on chromosomes 2 and 3 (Osorio-Guarín et al., 2020).

Although important economic diseases persist in cacao, wild populations exhibit disease resistance (Gutiérrez et al., 2016; Phillips-Mora et al., 2017). Only a few accessions have been used as a source of disease resistance to develop commercial clones, but more accessions should be studied for polygenic traits, such as disease resistance (Bailey et al., 2018; Gutiérrez et al., 2016). Meanwhile, using standard methods for artificial pod and leaf inoculation in germplasm collections and genotypes from different populations revealed new sources of disease resistance to FPR and BPR (Fister et al., 2020; Phillips-Mora et al., 2017; Puig et al., 2021). In addition, infection under field conditions also revealed new resistant genotypes. For example, the wild accession EBC-09 collected in the Colombian Amazon is resistant to FPR but not used commercially yet (Osorio-Guarín et al., 2020). Table 1 presents a summary of different candidate genes associated with disease resistance in cacao. Analysis of variation in these candidate genes in different genotypes can reveal potentially useful alleles for breeding programs. Thus, considering the continued demand for chocolate, breeding disease resistant varieties and renovating current cacao farms are more sustainable approaches than increasing
yield by planting new areas (Gutiérrez et al., 2016; Marelli et al., 2019; Phillips-Mora et al., 2012).

4.2 Genetic basis of quality traits

4.2.1 Chocolate flavor

A growing demand for chocolate also raises the need for related quality traits, including seed weight, fat content, low Cd content, and favorable flavor (CAOBISCO/ECA/FCC, 2015). The complex chocolate flavor depends not only on genotypes but also on soils, climate, farm management, bean fermentation, and roasting methods (Afoakwa et al., 2008; Kadow et al., 2013; Papalexandratou & Nielsen, 2016; Seguine & Meinhardt, 2014). Cacao seeds are broadly classified for fine aroma that has economic implications for chocolate products and the international cacao market (Afoakwa et al., 2008), where cacao with better aroma is higher in price compared to standard cacao. Only 12% of the global production is considered having a fine flavor. Fine chocolate flavor is commonly attributed to two genetic groups, Criollo and Nacional, and the hybrid Trinitarios (Criollo × Amelonado) (ICCO, 2022).

In the case of Nacional, the mature fruits are green due to the absence of anthocyanins and have a fruity–floral taste. Contrary, mature red fruits produced by standard cultivars such as CNN 5 are associated with unfavorable flavor (Motamayor et al., 2013). Based on the genome sequence of cultivar Matina1-6 (Amelonado group), association, and gene expression analysis, it was possible to identify polymorphisms in the candidate gene TcMYB113 responsible for fruit color variation (Motamayor et al., 2013). Thus, a single SNP associated with the absence of anthocyanins in mature green fruits allows to select for favorable flavor in breeding populations (Motamayor et al., 2013).

Fine chocolate flavor (floral and fruity) in Nacional is genetically controlled by different candidate genes distributed along the 10 cacao chromosomes (Colonges et al., 2021; Colonges,
Jimenez, et al., 2022). Two principal biosynthesis pathways are associated with floral taste, monoterpane biosynthesis, and the L-phenylalanine degradation (Colonges et al., 2021). In total, 27 candidate genes were associated with floral aroma in cacao, which were identified by GWAS studies and sensory analysis of 152 modern Nacional cultivars of Ecuador (Colonges et al., 2021). Furthermore, three additional biosynthesis pathways are involved in the synthesis of the fruity aroma of cacao: pyrazine production, sugar, and fatty acid (FA) degradation (Colonges, Jimenez, et al., 2022). In total, 327 candidate genes were associated with the fruity aroma in cacao (Colonges, Jimenez, et al., 2022).

Follow-up studies observed 1824 candidate genes determining cacao aroma in wild accessions of Ecuadorian Amazon (Colonges, Loor Solorzano, et al., 2022). Among these genes, 53 were also identified in previous studies investigating aroma traits in cultivated Nacional (Colonges, Loor Solorzano, et al., 2022). In addition, 81 candidate genes were identified as determinants of astringency, nonvolatile compounds, and cacao bitterness (Colonges, Seguine, et al., 2022). Overall, the balance among all these compounds provides the genetic basis for fine flavor formation in cacao and reveals a complex nature. Development of genetic markers based on candidate genes with high effects on favorable flavor would be necessary to specialized chocolate markets.

4.2.2 Cadmium (Cd) content

Cd is a heavy metal accumulated in edible parts of crops when transported from soils by the root system. The accumulation of Cd in food is neither a new problem nor unique to cacao. Fortunately, there is natural variation in Cd accumulation within crop species, and successful selection and breeding for low Cd-containing crops have been reported in barley, durum wheat, sunflower, rice, and soybean (Chen et al., 2007; Clemens et al., 2013; Grant et al., 2008; Lei et al., 2020).
The Cd transportation from soil to plant organs is associated with two main transporter genes encoding natural resistance-associated macrophage proteins and heavy metal ATPase (HMA), respectively (Clemens et al., 2013). For example, in rice (Oryza sativa L.), mutants with OsNRAMP5 defective function have successfully been used for breeding rice cultivars with low Cd accumulation and without affecting other agronomic traits (Ishikawa et al., 2012, 2019). Low Cd accumulation in rice was achieved by selecting varieties carrying functional alleles of OsHMA3, which promotes the Cd sequestration in root vacuoles, resulting in low Cd concentration in seeds (Ueno et al., 2010). Furthermore, an OsHMA3 allele with loss of function in Japonica rice cultivars was responsible for high Cd accumulation in grains (Yan et al., 2016). A transposable element presents in HvHMA3 allowed to enhance gene expression resulting in low Cd levels in barley grains (Lei et al., 2020). The low Cd allele originated from an international barley germplasm collection, being introduced into an elite cultivar by backcrossing, yielded grains with favorable agronomic traits, and low Cd accumulation (Lei et al., 2020). The introgressed barley lines yielded 0.1 mg Cd/kg compared to 0.4 mg Cd/kg in an elite cultivar; both groups of plants were grown in Cd contaminated soil (0.63 mg Cd/kg) (Lei et al., 2020).

Variation in cacao Cd accumulation called attention mainly because of international market restrictions for cacao powder and chocolate products with high Cd levels (between 0.10 and 0.80 mg Cd/kg) (The European Commission, 2014). The main factors related to the uptake of soil Cd by the cacao tree were soil pH, soil Cd availability, genotype, geographical location, and agronomic factors such as phosphate fertilizer (Oliveira et al., 2022). Different levels of Cd uptake have been observed among different cacao genotypes, suggesting genetic variation for seed Cd accumulation (Engbersen et al., 2019; Lewis et al., 2018). For example, in Trinidad and Tobago, 100 cacao accessions grown under the same conditions yielded different Cd contents in cacao seeds, ranging from 0.18 to 0.58 mg/kg for those classified as low Cd
accumulators, and from 1.72 to 2.16 mg/kg for high Cd accumulators (Lewis et al., 2018). Specifically, among the cacao genetic groups, the Contamana group exhibited high seed Cd content, whereas the Curaray group and hybrid populations such as Trinitario and Refractario are characterized by low Cd content (Lewis et al., 2018). Eleven cacao cultivars in Honduras demonstrated different Cd uptake rates (Engbersen et al., 2019). Accessions CAUCASIA-43, UF-676, and SPA-9 had high bean Cd accumulation (3–4 mg/kg), but UF 613 showed low bean Cd accumulation (∼1.3 mg/kg) (Engbersen et al., 2019). In cacao farms of Ecuador, the main factor affecting Cd accumulation was the presence of high Cd in soils combined with the natural ability of Cd uptake observed in Nacional genotypes and CCN 51 cultivars (Argüello et al., 2019). Overall, there is a tendency of cultivated cacao to increase seed Cd content when trees are grown in soils with high Cd content, which is common in Latin America (Meter et al., 2019; Vanderschueren et al., 2021). This accumulation pattern occurs despite environmental factors, probably because of a limited frequency of low Cd alleles within cacao cultivars.

The genes associated with Cd uptake in cacao have been studied recently. Expression and functional analysis of TcNRAMP5, TcHMA3, and TcHMA2 genes have demonstrated them as main Cd transporters in cacao (Moore et al., 2020; Ullah et al., 2018). To our knowledge, there is a gap between GWAS of Cd accumulation in seeds of cultivated cacao and genetic markers used for low Cd accumulation breeding. Cacao genotypes with low seed Cd accumulation (e.g., Red Amel 1/30, JA 5/39 and CL 27/50) can be tested as rootstocks for commercial clones to evaluate the genetic control of Cd accumulation (Lewis et al., 2018). However, soil–rootstock–scion interactions require further examination to validate potential low Cd cacao rootstocks (Fernández-Paz et al., 2021). For example, the commonly cultivated clone CCN 51 used as scion and the rootstock BN 34 accumulated higher Cd in root systems of juvenile cacao plants compared to seven different scion-rootstock combinations (Almeida et al., 2022).
4.2.3 Fat content and composition

The fat content of cacao seeds and their composition are vital for chocolate and cosmetics industries, and they demonstrated genetic and geographic variation (Almeida et al., 2008; Pires et al., 1998). Seed fat content varies among germplasm accessions from 45% to 60% in Bahia, Brazil (Pires et al., 1998), 51% to 58% in Rondônia, Brazil (Almeida et al., 2008), and 44% to 59% in Trinidad and Tobago (Khan et al., 2008). Cacao from different farms also varies for fat content with cacao cultivated in Ecuador showing a fat range of 45%–52%, whereas plantations in northeastern Peru show a low-fat content of 17%–30%. This variation was explained by geographic origin of samples and cultivated genotypes (Oliva-Cruz et al., 2021; Samaniego et al., 2021). On average, chocolate manufactures prefer a high fat content of 55%–58% in cacao seeds (CAOBISCO/ECA/FCC, 2015).

The fat profile of cacao is composed of three main FAs, which together account for more than 80% of the fat composition: saturated palmitic and stearic acids (C16:0, both with a 16-carbon chain, where the number before the colon specify the number of carbon atoms, and one after the colon, the number of double bonds) and monounsaturated palmitic oleic (C18:1). Other FAs are present in lower proportion, including saturated arachidic acid (C20:0) and the unsaturated linoleic acid (C18:2) (Mustiga et al., 2019; Samaniego et al., 2021; Torres-Moreno et al., 2015). This unique combination of FAs defines the texture, hardness, and an ideal melting point (37°C) of chocolate products at the mouthfeel (Afoakwa, 2016; CAOBISCO/ECA/FCC, 2015). A high ratio of stearic acid increases the hardness of cacao butter (Vieira et al., 2015), whereas higher levels of oleic acid produce soft cacao butter being less appropriate for chocolate manufacture (CAOBISCO/ECA/FCC, 2015; Vieira et al., 2015). Cacao yields soft butter when it grows in cooler climates (~24°C) revealing a strong effect of temperature on cacao butter profile (Vieira et al., 2015).
The fat profile of cacao also defines the nutritional characteristics of related products and is under genetic and environmental control. A healthier fat profile includes lower amounts of saturated and higher amounts of unsaturated fat, being preferred in chocolate products (Torres-Moreno et al., 2015). Specifically, a lower amount of palmitic acid in cacao fat is ideal because it reduces the risk of cardiovascular health problems, and a higher amount of stearic acid is favored because it is desaturated in the organism as oleic acid (Oliva-Cruz et al., 2021; Samaniego et al., 2021). Variations in fat profiles are observed in cacao populations. For example, the amount of palmitic acid is low in Criollo (20%–30%) but slightly higher in Nacional (26%–29%) and in accessions with Iquitos ancestry (26%–34%) (Mustiga et al., 2019; Samaniego et al., 2021). However, cacao grown at higher temperature (∼30–31°C) tends to produce a higher amount of palmitic acids, revealing a climate control (Mustiga et al., 2019). Furthermore, there is a negative correlation between total palmitic acid and the sum of stearic oleic and linoleic acids, which can be exploited to select genotypes with a healthier fat profile (Mustiga et al., 2019; Samaniego et al., 2021).

There are 84 genes orthologous to Arabidopsis that are potentially involved in fat production in cacao, of which 13 genes are unique to cacao (Argout et al., 2011). Specifically, the stearoyl-acyl carrier protein desaturase (SAD) gene family regulates fat synthesis and its composition in cacao (Li et al., 2019; Mustiga et al., 2019; Zhang et al., 2015). The activity of TcSAD defines stable levels of unsaturated and saturated FAs in cacao (Zhang et al., 2015). Among the eight putative isoforms of TcSAD, the expression of TcSAD1 was positively correlated with oleic acid and fat synthesis (Zhang et al., 2015). TcSAD7 is also involved in the synthesis and accumulation of oleic acid in cacao (Li et al., 2019).

Based on a mapping population of 420 progenies derived from the TSH 1188 × CCN 51 cross analyzed with a 6K SNP array and fat profiles from 3292 seed samples, it was possible to identify a significant QTL for a higher level in palmitic acid. The QTL marker explained a
4.5% increase in palmitic acid, which average content was 29% within the fat profile (Mustiga et al., 2019). This mapping population showed a low heritability for fat content (H2 = 0.14) mainly due to its complex nature and the effect of temperature and rainy seasons on seed fat production. However, it showed high heritability for palmitic acid (H2 = 0.43) and stearic acid (H2 = 0.38) (Mustiga et al., 2019), thus, demonstrating a good potential for breeding cacao genotypes with a healthier fat profile with lower palmitic and high stearic acids (Mustiga et al., 2019).

4.3 Genetic basis of agronomic traits

4.3.1 Seed traits and yield

The dry seeds of cacao are the valuable raw material for chocolate industries. Manufactures require uniform and heavier seeds (at least 1.0 g per fermented dried seed) because lighter and smaller cacao seeds have less fat percentage and more shell (CAOBISCO/ECA/FCC, 2015). Thus, the average cacao seed weigh is an important trait for breeding programs (Cilas et al., 2010; Doaré et al., 2020; Goenaga et al., 2009; Phillips-Mora et al., 2012). The seed weigh varies among genotypes. It also depends on the number of seeds per fruit, seed position within the fruit, and fruiting cycle (Doaré et al., 2020). Fruits with a high number of seeds (≥40 per fruit) tend to have lighter seeds (<0.94 g per seed) (Bekele et al., 2006). Seeds located in the apical sector of fruits also tend to be lighter, whereas longer fruiting cycles increase seed size and weight (Doaré et al., 2020).

Cacao populations show natural variation in seed weight. It ranged from 0.41 to 1.84 g per seed among 1900 accessions held in the International cacao collection of Trinidad (Bekele et al., 2020). Weight of wild cacao seeds ranged from 0.51 to 1.39 g compared to 0.55 to 1.88 g in cultivated clones (Bekele et al., 2006). In 11 Trinitario clones, even heavier seeds with a range of 1.15–1.84 g were identified (Bekele et al., 2020). In Brazil, seed weight followed a normal distribution across 221 clones and ranged from 0.3 to 2.8 g (Cilas et al., 2010).
addition, it was possible to structure accessions based on ancestry. Those with Trinitario and Criollo ancestry (commonly cultivated) have heavier seeds (>1.70 g) as a response to artificial selection. Contrary, wild populations showed more variation in seed weight (e.g., 0.90–1.45 g in the Guiana population) (Cilas et al., 2010). The seed weight showed a high heritability (H2 = 0.50), but the number of seeds per fruit had a lower heritability (H2 = 0.29). Thus, it is possible to breed cacao trees with high seed weight (>1.5 g) rather than increased number of seeds per fruit (>38) (Cilas et al., 2010).

Cacao candidate genes determining seed and yield traits were identified on chromosome IV via QTL mapping (Fernandes et al., 2020). Nine out of 13 candidate genes were involved in specialized functions such as sugar transport; other genes were involved in carbohydrate, lipid, and glucose metabolism (Fernandes et al., 2020). For each agronomic trait, such as number of pods (fruits) harvested per tree, dry seed weight, average yield per tree (kg of dry seeds/year), and pod index, it was possible to link at least one significant SNP contributing to the phenotypic variation of these traits, respectively (Fernandes et al., 2020). Pod index refers to the number of fruits needed to produce 1 kg of dry seeds; a low pod index (e.g., 21–25) is preferred because it helps to select trees that yield heavier and larger seeds (Bekele et al., 2020; Goenaga et al., 2009). A QTL with the highest effect explained 23.6% of the variation in the number of healthy pods harvested. This trait also showed a higher heritability (H2 = 0.75), thus, facilitating the selection of trees yielding more pods (Fernandes et al., 2020). Dry seed weight also exhibited a high heritability (H2 = 0.56), and among 16 QTL observed, one explained 14% of the variation for this trait (Fernandes et al., 2020).

4.3.2 Self-incompatibility (SI)

Self-incompatibility (SI) prevents self-fertilization in plants and helps to avoid inbreeding depression and to maintain plant genetic variability. It is common among crops (e.g., coffee, sunflower, and potato) and tropical trees, including cacao (Finkeldey & Hattemer, 2007; López
et al., 2021). Among cacao populations, SI is the predominant phenotype, but Amelonado and Criollo genetic groups exhibit self-compatibility (SC), and it is a trait of economic interest because SC trees are more productive (Lanaud et al., 2017; López et al., 2021; Royaert et al., 2011). Although cacao produces a lot of flowers (~125,000 annually per tree), no more than 10% of them are pollinated and less than 5% develop into mature fruits (Falque et al., 1995; Groeneveld et al., 2010). Thus, wild and planted cacao rely on midges flies from the Ceratopogonidae family for natural cross pollination (da Silva et al., 2016; Young, 1994). SI in cacao occurs when gametes fail to fuse in the ovary, preventing fertilization followed by flower abscission (Cope, 1962). Lanaud et al. (2017) provided a detailed description of the cacao SI system. Briefly, the complex SI system in cacao depends on different candidate genes that act independently and located on chromosome 1 (Chr1) and chromosome 4 (Chr4). Loci on both chromosomes were associated with gametic selection (Lanaud et al., 2017). Chr1 acts prior to gamete fusion and includes three candidate genes. For two of them expression was observed in style tissues. Chr4 includes nine candidate genes identified with SI reactions, including fruit drop (Lanaud et al., 2017).

Because cacao SI limits yield, artificial (handmade) pollination within and among cultivars help to identify and interplant compatible trees in the field to improve fruit productivity (Akoa et al., 2021; Forbes et al., 2019; López et al., 2021). However, artificial pollination has drawbacks because it can be influenced by rain, wind, and insect attacks and requires mature blooming trees (da Silva et al., 2016). Another alternative is to select SC cacao seedlings at an early stage using genetic markers (da Silva et al., 2016; Lanaud et al., 2017; Royaert et al., 2011). For example, seven SNPs markers associated with SC were discovered based on GWAS analysis (da Silva et al., 2016). The study used 5301 SNPs distributed over 10 cacao chromosomes, 291 diverse cacao trees, and flower set as phenotype (da Silva et al., 2016). The seven SNPs were located in the proximal end of CH4, where at least two genes were associated
with pollen rupture (da Silva et al., 2016). In addition, Lanaud et al. (2017) developed two SSR markers (mSI_303 and mSI_460) located in CH4 for predicting SC in cacao breeding populations.

Overall, planting SC trees, whose selection can be assisted using genetic markers (e.g., SNPs or SSRs), will reduce extra cost of pollination labor, and commercial cacao farms will not depend on specific designs where compatible clones need to be interplanted.

5 GENOMIC RESOURCES AVAILABLE FOR CACAO BREEDING

A better understanding of the genes contributing to agronomic traits in cacao was obtained from the sequence and annotation of two reference genomes, Criollo B97-61 (Argout et al., 2011, 2017) and Matina1-6 (Motamayor et al., 2013). From annotation of these genomes, it was possible to identify key genes involved in quality traits (e.g., fat biosynthesis, fine flavor, and fruit color). Details of Criollo and Matina cacao genomes are presented by Figueira and Camargo-Scotton (2020). These genomes are highly homozygous and do not represent wild and high yield cultivated hybrids (Morrissey et al., 2019). Thus, Pound 7, a wild cacao tree widely used in breeding because its high yield and resistance to BPD was sequenced using the low-cost MinION sequencer (Morrissey et al., 2019). Applying this sequencing strategy to cultivars of high local adaptability, for example, CCN 51 (Boza et al., 2014; Jaimez et al., 2022), can guide breeding and gene editing in specific traits like reduced bitterness and undesirable flavor. Table 2 presents the genomic statistics of these three genomes.

<table>
<thead>
<tr>
<th>Cacao genome</th>
<th>Estimated total genome size, Mbp</th>
<th>Total assembly, Mbp</th>
<th>Number of contigs</th>
<th>Number of scaffolds</th>
<th>Scaffold N50, Mbp</th>
<th>Number of genes</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Criollo B97-61 V2</td>
<td>430</td>
<td>326</td>
<td>7743</td>
<td>431</td>
<td>36.4</td>
<td>28,798</td>
<td>Argout et al., 2017</td>
</tr>
<tr>
<td>Matina1-6</td>
<td>445</td>
<td>331</td>
<td>20,103</td>
<td>714</td>
<td>34.4</td>
<td>29,408</td>
<td>Motamayor et al., 2013</td>
</tr>
<tr>
<td>Pound 7 (minimap)</td>
<td>442</td>
<td>393</td>
<td>2720</td>
<td>506</td>
<td>37.5</td>
<td>24,286</td>
<td>Morrissey et al., 2019</td>
</tr>
</tbody>
</table>

Abbreviation: Mbp, million base pairs.
Advances in cacao biotechnology, such as tissue culture, genetic transformation, and gene editing, offer new tools that enable precise breeding in cacao (Figueira & Camargo-Scotton, 2020; Wickramasuriya & Dunwell, 2018). For example, a pioneer study proves that CRISPR can be used in cacao, and improved resistance to BPD was achieved by knockout of the TCNPR3 gene (Fister et al., 2018). New experiments are needed to test if CRISPR is effective to improve yield, considering the co-localization of candidate genes with three SNP markers significantly associated with pod index and seed size (Bekele et al., 2022). In addition, physiological traits for genetic editing include small plant architecture and the abscission of mature fruits (Figueira & Camargo-Scotton, 2020). Crown reduction and semi-dwarfism reduce pruning intervals (Pereira et al., 2017), and abscission of mature fruits facilitates harvesting (Kuhn et al., 2010). Both traits reduce management costs, but studies on the genetic basis and candidate genes for these two traits are missing.

The Infinium cacao chip containing 6K SNPs (Livingstone et al., 2015) and the upgraded 15K SNP array (Livingstone et al., 2017) have been used in cacao GWAS and QTL analysis for improving disease resistance (Fernandes et al., 2018; Gutiérrez, Puig, et al., 2021; Romero-Navarro et al., 2017; Royaert et al., 2016) and yield (Fernandes et al., 2020; Romero-Navarro et al., 2017). Another application of genome-wide SNP data includes GS for predicting disease resistance and yield in breeding populations (McElroy et al., 2018). In addition, 3412 SNPs, selected from the 6 and 15K chocolate SNP chip, were used to develop cacao inbred lines aiming to generate hybrids and exploit potential heterosis in cacao breeding (Lopes et al., 2022). Specifically, the genome-wide SNP data helped to select cacao clones with up to 90% homozygosity within a germplasm collection in Brazil, and three generations of selfing monitored with SNP data can increase homozygosity up to 95% in clones like CCN 51 (Lopes et al., 2022).
Standard SNPs for cacao germplasm management aim to improve the efficiency and diversity of accessions maintained in *ex situ* collections (Gutiérrez, Martinez, et al., 2021; Mahabir et al., 2020). For example, SNP genotyping was used for validating cacao germplasm identity and detection of off-types in breeding programs (DuVal et al., 2017; Olasupo et al., 2018; Wang et al., 2020).

6 CONCLUSIONS AND OUTLOOK

Cacao agroforestry provides long-term land use with increased forest cover that benefits rural economies in the tropics. In addition, planting genetically improved cacao reduces the pressure on new land to satisfy the international demand of chocolate. We propose here research areas on germplasm management and agronomic traits that will mostly contribute to breeding new varieties required for sustainable cacao cultivation.

6.1 Conservation and use of cacao genetic diversity

In this review, the discussed geographic distribution of wild cacao in Amazonia represents trees mostly observed and/or collected along main rivers, suggesting that large gaps exist in the distribution presented in available studies (Figure 1). Considering the complexity of the Amazon basin, there is room for further plant collections and search for new cacao populations and germplasm. Unfortunately, the progressive deforestation of Amazonia is eroding the natural habitat not only of cacao populations but also of many more native plant species with crop and economic potential (Coelho et al., 2021; Neves & Heckenberger, 2019; Schultes, 1979). Thus, more genetic studies are urgently needed to study new cacao populations using next generation high-throughput sequencing and genotyping. For instance, to improve efficiency of cacao conservation, a 219 SNP genotyping panel was recently developed (Gutiérrez, Martinez, et al., 2021) providing an efficient tool to study and manage the genetic diversity of wild and admixed accessions, including those that are maintained in around 47
cacao germplasm collections. Genetic gaps, off-types, and redundancy can be identified through SNP genotyping. Moreover, specific SNP profiles can identify commercial clones and parent trees carrying valuable agronomic traits in breeding programs.

Continued characterization of agronomic traits in different genetic groups (e.g., through genotyping accessions maintained in germplasm collections) can reveal new alleles valuable for breeding agronomic traits. It is important to note that multiyear agronomic characterization requires standard protocols shared among breeding programs for phenotyping traits, such as yield (e.g., pod index), seed weight, drought, and disease resistance. For example, new available methods can be adapted by different cacao research programs to evaluate yield and disease resistance (Jaimez et al., 2020), resistance to BPR in seedlings (Delgadillo-Durán et al., 2020), and on detached fruits (Puig et al., 2021). In addition, standard protocols to evaluate cacao quality and flavor are available (ISCQF, 2020). Overall, standard phenotyping is a priority that allows replicability and comparability of trait data among cacao breeding programs.

6.2 Disease resistance

Characterization of cacao resistance to FPR can benefit from the identified centers of genetic diversity of M. roreri in Magdalena valley of Colombia, Coastal Ecuador, and Ecuadorian Amazon. Thus, different cacao genetic groups can be tested against different M. roreri isolates, potentially identifying new sources of disease resistance. In addition, studying wild populations of M. perniciosa in Amazonia (e.g., Colombia, Peru, and Ecuador) can provide more information about genetic diversity, dispersals patterns, and genotypes of M. perniciosa causing WBD in cultivated cacao. Moreover, it will help to spot forest areas in Amazonia for collecting wild cacao that likely evolved resistance to WBD and BPR.
6.3 Climate adaptation

Breeding and planting cacao trees tolerant to drought and heat in West Africa can reduce the risk of deforestation associated with new suitable planting areas. Different cacao populations can be tested in controlled drought experiments aided by GWAS analysis to identify potential SNP markers in root phenotypes (e.g., angle and biomass) that improve water use efficiency. Of particular interest are the populations that originated along the border of Peru and Ecuador, and in the south of Peru, where wild trees grow naturally with precipitations around 1000 mm/year (Figure 2), potentially providing a natural source of germplasm to exploit drought tolerance traits.

6.3 Low seed cadmium (Cd) accumulation

Two main genes have been identified as main Cd transporters from soil to the cacao tree—TcNRAMP5 and TcHMA (Moore et al., 2020; Ullah et al., 2018). In addition, there is evidence of natural variation in seed Cd accumulation among cacao genotypes growing under the same soil conditions (Lewis et al., 2018). However, we neither know whether TcNRAMP5 and TcHMA gene variants are correlated with seed Cd accumulation levels, nor how these potential variants are distributed in cacao populations. By identifying gene variants (e.g., using SNP markers) associated with low and high seed Cd accumulation, breeders can develop new varieties to limit the Cd exposure on chocolate consumers.

6.5 Gene editing

Natural tree populations provide valuable genetic diversity for breeding programs. Using this variation and new genomic technologies, such as gene editing, it will be possible to accelerate tree genetic improvement (Cao et al., 2022). For example, the efficiency of gene editing using the CRISPR/Cas9 method was tested in cacao accession PSU SCA6 by knocking out the TcNPR3 gene that resulted in enhanced resistance to Phytophthora tropicalis (Fister et al.,
Elite cacao clones can directly be improved by CRISPR-based gene editing, but it requires high quality reference genomes for optimizing specific gRNA constructs and to avoid off-target products (Morrissey et al., 2019). In addition, cacao tissue culture needs to be optimized, although protocols for somatic embryogenesis using staminode explants were adapted in nine cacao cultivars (Jones et al., 2022). In addition, further gene editing studies can validate candidate gene functions (e.g., loss of the function mutations) in important biological processes of cacao, such as disease resistance, and SC and Cd uptake. For example, functionally characterized Cd uptake causative genes (TcNRAMP5 and TcHMA) could be further validated in wild populations and clones. However, the potential of CRISPR-based technology is limited to agronomic traits controlled by a few major genes, genome editing of polygenic complex traits is a complicated task that may cause unintended results (Cao et al., 2022). Ultimately, the acceptance of gene editing and genetic modifications of cacao will depend on public acceptance and official regulations, but farmers and consumers will directly benefit from improved cacao.

6.6 Genomic selection (GS)

GS is more often implemented now in tree improvement due to the high accuracy of prediction of complex phenotypes that shortens breeding cycles (Grattapaglia, 2022). In cacao, GS using a 15K SNP genotyping array proved to be effective in improving quantitative traits, such as disease resistance and yield (Romero et al., 2017; McElroy et al., 2018). High-density SNP genotyping arrays can provide long-term benefits like higher genetic gain and better predictions (Grattapaglia, 2022). For example, medium to high-density SNP genotyping arrays are common in breeding tropical perennial crops, such as Eucalyptus (60K SNPs) and oil palm (200K SNPs) (Seyum et al., 2022). In addition, experimental approaches such as development of cacao inbred lines as elite parents can be used for mate selection and exploitation of hybrid vigor (Lopes et al., 2022), followed by GS of untested hybrids across different environments.
Methods and practical guidelines for designing GS programs specifically for cacao are much needed similar to other tree crops (see reviews by Grattapaglia, 2017, and Isik, 2022). The private sector and research agencies collaborated to test cacao hybrids and shorten the breeding cycle of disease resistance and yield under GS schemes (Romero et al., 2017; McElroy et al., 2018). However, challenges still remain, such as building a mature breeding population (e.g., by eliminating low yield and susceptible individuals at an early stage), high cost of genotyping, multiyear data collection and complexity of phenotyping disease resistance in training populations, and testing for $G \times E$ interactions in locations with higher disease pressure.

**AUTHOR CONTRIBUTIONS**

Helmuth E. Nieves-Orduña: Conceptualization; Funding acquisition; Investigation; Visualization; Writing — Original Draft; Writing — Review & Editing. Konstantin V. Krutovsky: Conceptualization; Investigation; Supervision; Validation; Writing — Review & Editing. Oliver Gailing: Conceptualization; Funding acquisition; Investigation; Project administration; Supervision; Validation; Writing — Review & Editing.

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CONFLICT OF INTEREST STATEMENT

The authors declare no conflict of interest.
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Paper 2

Geographic patterns of genetic variation among cacao (*Theobroma cacao* L.) populations based on chloroplast markers

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Geographic Patterns of Genetic Variation among Cacao (*Theobroma cacao* L.) Populations Based on Chloroplast Markers

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**Keywords:** cacao; chloroplast haplotypes; geographic origin; chocolate; crop dispersal; SSR; microsatellite markers
Abstract

The cacao tree (*Theobroma cacao* L.) is native to the Amazon basin and widely cultivated in the tropics to produce seeds, the valuable raw material for the chocolate industry. Conservation of cacao genetic resources and their availability for breeding and production programs are vital for securing cacao supply. However, relatively little is still known about the phylogeographic structure of natural cacao populations. We studied the geographic distribution of cpDNA variation in different populations representing natural cacao stands, cacao farms in Ecuador, and breeding populations. We used six earlier published cacao chloroplast microsatellite markers to genotype 233 cacao samples. In total, 23 chloroplast haplotypes were identified. The highest variation of haplotypes was observed in western Amazonia including geographically restricted haplotypes. Two observed haplotypes were widespread across the Amazon basin suggesting long distance seed dispersal from west to east in Amazonia. Most cacao genetic groups identified earlier using nuclear SSRs are associated with specific chloroplast haplotypes. A single haplotype was common in selections representing cacao plantations in west Ecuador and reference Trinitario accessions. Our results can be used to determine the chloroplast diversity of accessions and in combination with phenotypic assessments can help to select geographically distinctive varieties for cacao breeding programs.
1 INTRODUCTION

Cacao trees (*Theobroma cacao* L.) are native to the Amazon basin, and their valuable seeds are the raw material for the chocolate industry (Bartley, 2005; Motamayor et al., 2008). Cultivated by 5–6 million small-holder farmers in Latin America, Africa, and Asia, cacao crops are vitally important for local economies in these continents (Beg et al., 2017; Rice & Greenberg, 2000). As an important tropical crop, there is a constant need to develop high-yielding and disease-resistant varieties (Bennett, 2003; Goenaga et al., 2009; Phillips-Mora et al., 2013). Thus, conservation and use of cacao genetic diversity are essential not only for sustainable cultivation in producing countries, but also for diverse and growing needs of the chocolate industry and consumers (Bartley, 2005; CacaoNet, 2012; Laliberté et al., 2018; Zhang et al., 2011; Zhang & Motilal, 2016).

Western Amazonia is considered to be a putative center of origin of cacao based on high phenotypic (Bartley, 2005; Cheesman, 1944; Cuatrecasas, 1964) and genetic (Cornejo et al., 2018; Motamayor et al., 2008; Sereno et al., 2006) diversity. This part of the Amazon is also a region of high biodiversity (Bass et al., 2010; Garcia-Davila et al., 2020; Myers et al., 2000; Hans Ter Steege et al., 2003; Tuomisto et al., 1995) that in the past served as a forest refugium favoring accumulation of high tree diversity (Prance, 1982; van der Hammen & Hooghiemstra, 2000). Western Amazonia is also considered a center of crop domestication (Clement et al., 2010; Clement, 1999; Clement et al., 2015; Meyer et al., 2012). Examples of cultivated tree species, whose probable origin of domestication is western Amazonia, include fruit trees, such as the ice cream bean tree (*Inga edulis* Mart.), tree grape (*Pourouma cecropiifolia* Mart.), caimito (*Pouteria caimito* Radlk.), and the Amazon nut tree (*Bertholletia excelsa* Bonpl.) (Levis et al., 2017). At least 29 crop species and 38 utilizable palms with probable origin in western Amazonia highlight the importance of the plant genetic resources observed in this part of the Amazon (C. R. Clement, 1999; Paniagua-Zambrana et al., 2007).
Cacao populations are organized in 10 genetic groups, some of which are called varieties (Amelonado, Contamana, Criollo, Curaray, Guianna, Iquitos, Marañon, Nacional, Nanay, and Purús), spread over the native range in South America (Motamayor et al., 2008). This classification is based on the Bayesian cluster analysis of 96 nuclear SSR markers genotyped in 952 cacao accessions selected from ex situ collections (Motamayor et al., 2008).

The cultivated cacao was traditionally classified into Criollo and Forastero (an umbrella designation for the Amazonian populations) varieties and a Trinitario variety, a natural hybrid of Criollo × Forastero (Bartley, 2005; Cheesman, 1944). However, the contribution of some of the 10 cacao groups to cultivated cacao is relatively small, which is represented mainly by three varieties: Amelonado, Criollo, and Nacional (Bartley, 2005; Bennett, 2003; Cornejo et al., 2018; Zhang et al., 2011; Zhang & Motilal, 2016). However, during domestication, the Criollo variety accumulated a large proportion of high-frequency deleterious mutations that affect fitness (Cornejo et al., 2018), necessitating the use of more diverse material for cacao breeding (Bennett, 2003; Cornejo et al., 2018; Zhang et al., 2011; Zhang & Motilal, 2016).

One of the approaches to better understand genetic resources available for cacao breeding is to characterize historical patterns of seed dispersal and the origin of cultivated populations using chloroplast (cpDNA) markers, such as chloroplast microsatellites (cpSSRs) (Yang et al., 2011). They are polymorphic markers of the chloroplast genome and typically maternally inherited in angiosperms, including cacao, and propagated via seeds. Therefore, they are used preferentially for phylogeographic analysis (Lemes et al., 2010; Weising & Gardner, 1999). Nevertheless, a set of standard nuclear SSRs (Irish et al., 2014; Motilal et al., 2011; Saunders et al., 2004) and SNPs (Jemmy et al., 2014; Ji et al., 2013; Lukman et al., 2014; Padi et al., 2015) are preferred for cacao identification in germplasm collections and breeding experiments.
Specifically, chloroplast markers are useful to understand geographic variation and dispersal of tropical trees (Caron et al., 2000; Cavers et al., 2003; Dick, 2010; Hamilton, 1999; Lemes et al., 2010). For instance, these markers revealed long distance transfer of seeds for plantation establishment in *Dalbergia sissoo* Roxb. (Pandey et al., 2004) and helped to select specific haplotypes to improve site adaptation in cultivated *Pinus armandii* Franch. (Jia et al., 2020). In cacao, the historical transfer of reproductive material for cultivation has focused on genotypes exhibiting disease resistance and flavor traits (Bartley, 2005).

Nine specific chloroplast microsatellites (cpSSR) are available for cacao genetic analysis (Yang et al., 2011). In cacao farms and localities in Trinidad and Tobago, these markers allowed the identification of seven haplotypes among Trinitarian cacao populations (Yang et al., 2013). Maternal lineages (seeds) used for cultivar development in Trinidad apparently originated from cacao populations of Central America, Peru, and Venezuela (Yang et al., 2013).

Ten haplotypes were identified by sequencing the chloroplast intergenic *trnH-psbA* spacer region of cultivated cacao trees in Soconusco, southern Mexico (Gutiérrez-López et al., 2016), where cacao cultivation has been recorded since colonial times (Coe & Coe, 2013). Gutiérrez-López et al. (2016) confirmed the introduction of a few mother trees as founder material for Soconusco plantations.

Although these studies did not include a wide range of samples within the natural distribution of cacao populations in the Amazon basin, they validated the use of chloroplast markers for managing cacao genetic resources. Here, we show how chloroplast genetic diversity is geographically distributed in Amazonia, and how it is represented in common cacao genotypes by using cpSSR markers. We hypothesize a center of cpDNA diversity in western Amazonia, considering the high genetic diversity observed in western cacao populations and
the recognized origin of the species in this region. Our objectives were to: (i) study the geographic distribution of cpDNA variation in a sample of cacao trees collected in the Amazon basin, and in samples of cultivated cacao from west Ecuador, (ii) identify cpDNA variation in breeding populations, and (iii) determine the correspondence between cacao chloroplast and nuclear DNA variation.

2. MATERIALS AND METHODS

2.1. Plant Material

Fresh leaves of 233 individual cacao trees were obtained for DNA extraction from the living collection maintained by the International Cocoa Quarantine Centre, Reading University, UK. Leaf samples of 154 cacao trees were selected based on widespread geographic origin from Amazon forests in five countries (Figure 1) (31 locations and 1–28 individuals per location, Table S1). This material corresponds to seeds or budwood collected during different expeditions in 1938–1988, aiming to obtain plant material carrying disease resistance traits and to study the species’ ecology and variability (Allen, 1988; Pound, 1938; Sallée, 1987; Turnbull & Hadley, 2019; Vello & Madeiros, 1965). Although these plant expeditions took place in Amazonian forests, some collection sites possibly experienced pre-Columbian silvicultural management like ancient cacao cultivation or translocation (Barlow et al., 2012; C. R. Clement et al., 2015; Peters, 2000).

Samples of 30 individuals of cultivated cacao originated from nine different cacao farms in the coastal valley of Ecuador (1–6 individuals per location, Table S2, Figure 1). This group of germplasm is known as Refractario cacao and includes the progeny of approximately 80 different trees selected in 1937 for their resistance to witches’ broom disease (Pound, 1938, 1943; Turnbull & Hadley, 2019; Zhang et al., 2008).
Additionally, leaf samples of 49 clones used for plantations and breeding programs were analyzed. They included the following genotypes: UF (United Fruit); ICS (Imperial Collection Selection); CRU (Cocoa Research Unit); TSH (Trinidad Selected); EET (Estación Experimental Tropical); and VB (Vassoura de Bruxa). The complete list of genotypes and their breeding origin in ten countries from Latin America and the Caribbean are presented in Supplementary Table S3.

Among the 233 samples analyzed, 83 different clones were also included in the analysis of [2] and represented different cacao genetic groups with different geographic origin: Amelonado (8), Contamana (6), Curaray (7), Guiana (2), Iquitos (23), Maraño (23), Nacional (2), and Nanay (12) (Tables S1 and S3). The International Cocoa Germplasm Database (ICGD) (Turnbull & Hadley, 2019) provides details about agronomic traits, breeding programs, plant collection expeditions and geographic origin of the 234 accessions analyzed.
2.2. DNA Extraction

To extract DNA, we used about 1 cm² of leaf tissue and the DNeasy 96 Plant Kit (Qiagen, Hilden, Germany). DNA was diluted 1:10 before PCR amplification.

2.3. Chloroplast DNA Markers

We tested the amplification and diversity of ten universal chloroplast markers (ccmps) (Weising & Gardner, 1999) in eight samples from distant regions. Additionally, we tested nine chloroplast microsatellite markers developed earlier for T. cacao (CaCrSSRs) based on the cacao chloroplast genome (Yang et al., 2011). Monomorphic amplification products were obtained with all ccmp markers, but six out of the nine cacao chloroplast markers showed clear polymorphisms, which were used then to screen the 235 samples. Four of them (CaCr2, CaCr4, CaCr5, and CaCr8) represented mononucleotide, one (CaCr1)—pentanucleotide, and one—octonucleotide (CaCr9) repeats. Although it is difficult to genotype mononucleotide repeats, it is easier to do it for chloroplast SSRs than for nuclear SSRs because they are haploid and their fragment size can be easier determined due to the lack of interference with another allele such as in diploid nuclear SSRs. Different alleles were also verified by running respective samples side-by-side during the same electrophoretic run. When the allele calling was no clear, we did repetitions to ensure a correct allele scoring.

The following six polymorphic chloroplast DNA (cpDNA) markers were amplified by PCR in two multiplexes: (1) CaCrSSR1, CaCrSSR2, and CaCrSSR4, and (2) CaCrSSR5, CaCrSSR8, and CaCrSSR9. A M13 tail (5’-CACGACGTTGTAAACGAC-3’) and a PIG tail (5’-GTTTCTT-3’) were attached to the 5’ ends of forward and reverse primers, respectively (Kubisiak et al., 2013).

The PCR reaction mix for each primer in 14 µL volume contained: 1 µL of genomic DNA (about 0.6 ng/µL), 5.7 µL ddH2O; 1.5 µL PCR buffer (10× Buffer B1 from Solis BioDyne,
containing Tris–HCl and (NH₄)₂SO₄), 1.5 µL MgCl₂ (25 mM), 1 µL dNTP (2.5 mM of each dNTP), 0.2 µL (5 U/µL) HOT FIREPol® Taq Polymerase from Solis BioDyne, and 0.2 µL of each forward primer, 0.5 µL of each reverse primer, and 1 µL of the M13 primer (6-FAM) for the first multiplex, but 0.3 µL and 0.75 µL of the forward and reverse primers for the CaCrSSR5 marker, respectively, 0.1 µL and 0.25 µL of the forward and reverse primers for the CaCrSSR8 marker, respectively, 0.2 µL and 0.5 µL of the forward and reverse primers for the CaCrSSR9 marker, respectively, and 1 µL M13 primer (HEX) for the second multiplex. Concentration of all primers was 5 pM/µL.

The PCR conditions for both multiplexes were 95 °C for 15 min followed by 35 cycles of 95 °C for 1 min, 50 °C for 1 min, 72 °C for 1 min, followed by final 72 °C for 20 min, and a hold at 16 °C.

By using 1.5% agarose gel electrophoresis single bands for all PCR reactions were visualized, and the dilution ratio for the PCR product was determined. PCR products diluted at 1:10 were run on an ABI 3130xl Genetic Analyser (Applied Biosystems, Foster City, CA, USA) with GS 500 ROX used as internal size standard.

2.4. Data Analysis

Allele size analysis was performed using GeneMapper version 4.1 (Applied Biosystems, Foster City, CA, USA). Haplotypes based on all six chloroplast markers and their frequencies were determined using the Haplotype Analysis version 1.05 software (Eliades & Eliades, 2009). This software was also used to estimate total (HT) and within population (HS) haplotypic diversity, genetic differentiation (FST) between western and eastern natural populations, and between natural populations and cultivated cacao in Ecuador. The haplotype network to visualize the relationships among haplotypes was generated using the Network 5.0.1.1 software (Bandelt et al., 1999). First, a “rdf” file was created in GenAlEx 6.5 (Peakall & Smouse, 2012), and then
it was reformatted and saved in Network 5.0.1.1 as an “ych” file and used by this software as input file to generate the haplotype network using the median joining method. The created output file (“out”) was saved and used for visualization (Bandelt et al., 1999).

3. RESULTS

3.1. Identification of Haplotypes

In total, 26 alleles were identified for the six cacao chloroplast markers in 233 samples. The CaCrSSR1 marker with a pentanucleotide repeat was the most polymorphic marker with nine alleles. Six alleles were observed at CaCrSSR5, five at CaCrSSR2, three at CaCrSSR4, and two at each CaCrSSR8 and CaCrSSR9. These alleles allowed us to detect 23 haplotypes. Table 1 shows the allele compositions of these haplotypes and their frequencies. Six samples were excluded from the data analysis due to their incomplete genotyping (Tables S1–S3).

**TABLE 1. Theobroma cacao L. chloroplast haplotypes and their frequencies observed in different locations in the Amazon, plantations, and breeding populations.**

<table>
<thead>
<tr>
<th>Chloroplast haplotype</th>
<th>CaCr1</th>
<th>CaCr2</th>
<th>CaCr4</th>
<th>CaCr5</th>
<th>CaCr8</th>
<th>CaCr9</th>
<th>Western Amazonia</th>
<th>Other locations*</th>
<th>Plantations</th>
<th>Breeding populations</th>
<th>N</th>
<th>Frequency, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>361</td>
<td>243</td>
<td>177</td>
<td>207</td>
<td>309</td>
<td>345</td>
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<td>2</td>
<td>2</td>
<td>4</td>
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<td>2.9</td>
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<td>1</td>
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<td>1</td>
<td>4</td>
<td>1</td>
<td>0.4</td>
</tr>
</tbody>
</table>
3.2. Western Amazonia, a Center of Haplotype Diversity

We observed high cpDNA genetic diversity in western Amazonia, with 19 haplotypes detected in this region in total (Figure 2 and Figure 3). In Peruvian Amazon, 11 haplotypes were observed in three river systems, with an average of 4.6 haplotypes. The highest number of haplotypes was found along the Amazon River at Iquitos, followed by the Marañon and Nanay rivers with haplotypes H22, H16, and H19 being the most frequent, respectively. In addition, five haplotypes were observed in the south and central part of Peru along the Urubamba and Ucayali rivers (Figure 2). Nine haplotypes were observed in Ecuadorian Amazon along the Coca and Napo rivers, with haplotypes H9 and H5 being the most frequent, respectively. Finally, two haplotypes, H22 and H12, were observed in the Colombian Amazon (Figure 3). A high total haplotypic diversity (HT = 0.725) and a large number of haplotypes (Nh = 19) were detected in western Amazonia in 28 locations (Figure 2 and Table 2). The within population haplotypic diversity was also high (HS = 0.676) in this area.

<table>
<thead>
<tr>
<th>Amazonia</th>
<th>N</th>
<th>NLoc</th>
<th>Nh</th>
<th>Hs</th>
<th>HT</th>
<th>FST</th>
</tr>
</thead>
<tbody>
<tr>
<td>Western</td>
<td>115</td>
<td>28</td>
<td>19</td>
<td>0.676</td>
<td>0.725</td>
<td>0.068</td>
</tr>
<tr>
<td>Eastern</td>
<td>32</td>
<td>3</td>
<td>2</td>
<td>0.037</td>
<td>0.087</td>
<td>0.572</td>
</tr>
</tbody>
</table>

N - sample size, NLoc - number of locations, Nh - number of haplotypes, Hs - within population haplotypic diversity, HT - total haplotypic diversity, FST - differentiation among populations within regions.

In contrast, only two haplotypes (H16 and H17) were found in central and eastern Amazonia (Figure 2). Both the total (HT = 0.087) and the within population (HS = 0.037) haplotypic diversity were low in eastern Amazonia (Table 2). When compared, eastern and western cacao populations were highly divergent (FST = 0.500), since they shared only two haplotypes in this study (Figure 2).
FIGURE 2. Distribution and frequencies of chloroplast haplotypes among *Theobroma cacao* L. populations in the Amazon basin. The rectangle represents the area of high haplotype diversity in northwestern Amazon (displayed in detail in Figure 3). The green dots represent samples from natural populations, the blue dots plantations, and the red dots accessions in breeding stations (CATIE-Tropical Agricultural Research and Higher Education Center, ICGT-International Cocoa Genebank Trinidad). The size of the circles is proportional to the number of samples per location.

FIGURE 3. Distribution and frequencies of chloroplast haplotypes among *Theobroma cacao* L. populations in western Amazonia. The green dots represent samples from natural populations, the blue dots plantations, and the red dots accessions in breeding stations. The size of the circles is proportional to the number of samples per location.
3.3. Geographically Restricted Haplotypes in Western Amazonia

In Peru, seven geographically restricted haplotypes occurred at three locations: the confluence of the Marañon and Amazon rivers (H3, H14, H15, H18, H23), Morona River (H2) and Napo River (H11). In addition, haplotype H1 was locally common in the south of Peru along the Urubamba-Ucayali River (Figure 2).

In Ecuador, the geographically restricted haplotype H13 was observed along the Napo River. Moreover, haplotypes H5 and H9 were common in different locations along the Napo basin at the border with Colombia, and haplotype H5 was common along Upano River. Haplotype 12 was observed in the Colombian Amazon at Caquetá River (Figure 3).

3.4. Correspondence between Chloroplast and Nuclear DNA Variation

There is an agreement between the chloroplast haplotypes observed in 83 individuals (Tables S1 and S3) included in Motamayor et al. (2008) and eight cacao genetic groups. Each group generally has one dominant haplotype and other related haplotypes. Results may suggest that the cacao genetic groups proposed by Motamayor et al. (2008) are heterogeneous in terms of the chloroplast variation (Table 3). However, to study in detail patterns of correspondence between cpDNA and nuclear variation in cacao, sampling for cpDNA analysis should ideally include all individuals analyzed by Motamayor et al. (2008).

The dominant haplotypes within the genetic groups in the Peruvian Amazon were H22 in Iquitos, H19 in Nanay, H16 in Marañon, and H20 in Contamana in central Peru. In the Ecuadorian Amazon the dominant haplotype was H9 in Curaray and H10 in Nacional, and in central-eastern Amazonia H17 in Amelonado and H16 in Guiana (Table 3).
TABLE 3. Correspondence between nuclear and chloroplast DNA analyses in cacao (*Theobroma cacao* L.). Cacao genetic groups, haplotype frequency and number of samples analyzed in this study and in Motamayor et al. (2008).

<table>
<thead>
<tr>
<th>Genetic group</th>
<th>Chloroplast haplotype (F)</th>
<th>N (this study)</th>
<th>N (Motamayor et al. 2008)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nanay</td>
<td>19 (0.91), 3 (0.08)</td>
<td>12</td>
<td>121</td>
</tr>
<tr>
<td>Iquitos</td>
<td>22 (0.86), 23 (0.04), 20 (0.04), 16 (0.04)</td>
<td>22</td>
<td>75</td>
</tr>
<tr>
<td>Maraño</td>
<td>16 (0.78), 14 (0.08), 17 (0.04), 15 (0.04), 14 (0.04)</td>
<td>23</td>
<td>130</td>
</tr>
<tr>
<td>Guiana</td>
<td>16 (1.0)</td>
<td>2</td>
<td>51</td>
</tr>
<tr>
<td>Amelonado</td>
<td>17 (0.62), 22 (0.12), 16 (0.12), 5 (0.12)</td>
<td>8</td>
<td>63</td>
</tr>
<tr>
<td>Contamana</td>
<td>20 (0.51), 1 (0.33), 19 (0.16)</td>
<td>6</td>
<td>59</td>
</tr>
<tr>
<td>Curaray</td>
<td>9 (0.33), 2 (0.33), 13 (0.16), 20 (0.16)</td>
<td>6</td>
<td>87</td>
</tr>
<tr>
<td>Nacional</td>
<td>10 (0.50), 6 (0.50)</td>
<td>2</td>
<td>36</td>
</tr>
</tbody>
</table>

Four haplotypes were common within nuclear genetic groups: haplotype H16 was observed from west to east in the Amazon basin in Maraño (78%), Amelonado (12%), Iquitos (4%) and Guiana (100%). It is worth noticing that Maraño, Guiana, and Amelonado were also related groups based on nSSRs (Figure 2 in Motamayor et al. 2008). Haplotype H17 occurred together with the closely related H16 in Maraño demonstrating a phylogeographic pattern (Figure 4). Finally, haplotype H19 was shared between distinct clusters Nanay (91%) and Contamana (16%), and haplotype H20 was detected in the distantly related groups Contamana (50%) and Curaray (16%) (Table 3, Figure 2 in Motamayor et al. 2008).

3.5. Haplotypes in Ecuadorian Plantations

Eight haplotypes were identified in 29 samples in nine cacao plantations along the coast of Ecuador, with haplotype H10 being the most frequent one (72%) from north to southwest Ecuador (Figure 3). The geographic origin of H10 is associated with three locations in four
samples: Napo basin in Ecuador (N = 2), Morona (N = 1) and Marañón (N = 1) rivers in Peru (Figure 3).

**FIGURE 4.** Network of chloroplast haplotypes (1–23) in *Theobroma cacao* L. populations. Each circle represents a specific haplotype, and its size reflects the respective haplotype frequency. Vertical bars in the lines connecting haplotypes indicate hypothetical mutations separating haplotypes. The distance between neighboring vertical lines is the same, but it was condensed in some lineages for better space use and visualization of the network.

Cacao farms in Ecuador also contained haplotype H2 being the second most frequent haplotype in plantations (Table 1, Figure 3). In three locations totaling three samples, H2 was observed along Napo River in Ecuador and Morona River in Peru. However, H2 and H10 may also occur in other Amazonian cacao populations that were not analyzed here.

H4 and H8 were rare haplotypes in Ecuadorian plantations observed only in Hacienda Balao and Vuelta Larga, respectively. These haplotypes were not observed in any natural population studied here (Figure 2 and Figure 3).
Low total ($HT = 0.116$) and within population ($Hs = 0.075$) haplotypic diversity were observed in plantations of western Ecuador due to the high frequency of H10 and relatively low sample size in plantations ($N = 29$). Differentiation between cultivated and samples collected in Amazonian forest was high ($F_{ST} = 0.500$).

### 3.6. Haplotypes Observed in Breeding Populations and Cultivars

Cacao breeding has been mostly relying on vegetatively propagated accessions, such as Imperial College Selection (ICS), Iquitos Mixed Calabacillo (IMC), Nanay (NA), Parinari (PA), Pound, Scavina (SCA), and United Fruit (UF) (Bekele & Phillips-Mora, 2019; L.A.S. Dias, 2001; Lopes et al., 2011). Table 4 presents haplotypes detected in reference accessions and their potential geographic sources. For example, SCA 6, Pound 18, PA 139, and IMC 67 showed H5, H16, H20, and H22 haplotypes, respectively, and the geographic origin of these accessions ranged from central to northern Peru (Figure 2). In addition, cultivars, such as UF 273 (Nacional × Amelonado), ICS, and EET (Nacional × Unknown) revealed the common haplotype H10 observed in Ecuadorian plantations (Figure 3).

**TABLE 4.** Chloroplast haplotypes observed in cacao (*Theobroma cacao* L.) breeding populations, tentative geographic origin of haplotypes, source, and breeding program.

<table>
<thead>
<tr>
<th>Clone name</th>
<th>Source</th>
<th>Haplotype</th>
<th>Tentative geographic origin</th>
<th>Breeding program origin</th>
</tr>
</thead>
<tbody>
<tr>
<td>CC 137</td>
<td>Cacao Center</td>
<td>2</td>
<td>Morona River, Peru</td>
<td>CATIE</td>
</tr>
<tr>
<td><em>UF 168</em></td>
<td>United Fruit selections</td>
<td>3</td>
<td>Nanay River, Peru</td>
<td>UF Company, Costa Rica</td>
</tr>
<tr>
<td>NA 399</td>
<td>Nanay</td>
<td>3</td>
<td>Nanay River, Peru</td>
<td>ICG,T</td>
</tr>
<tr>
<td>ICS 68</td>
<td>Imperial Collection Selection</td>
<td>5</td>
<td>Western Amazonia Iquitos, Peru</td>
<td>ICG,T</td>
</tr>
<tr>
<td>POUND 18</td>
<td>Pound Collections</td>
<td>5</td>
<td>Western Amazonia Iquitos, Peru</td>
<td>ICG,T</td>
</tr>
<tr>
<td>CC 252</td>
<td>Cacao Center</td>
<td>9</td>
<td>The northeast of Ecuador (Coca and San Miguel rivers)</td>
<td>CATIE</td>
</tr>
<tr>
<td><em>UF 712</em></td>
<td>Programa Mejoramiento de Cultivos Tropicales</td>
<td>9</td>
<td>The northeast of Ecuador (Coca and San Miguel rivers)</td>
<td>CATIE</td>
</tr>
<tr>
<td><em>UF 273</em></td>
<td>United Fruit selections</td>
<td>10</td>
<td>The northeast of Ecuador (Coca River and north of Peru (Morona and Marañon rivers))</td>
<td>UF Company, Costa Rica</td>
</tr>
<tr>
<td><em>ICS 5, 15, 42, 48, 63</em></td>
<td>Imperial Collection Selection</td>
<td>10</td>
<td>The northeast of Ecuador (Coca River and north of Peru (Morona and Marañon rivers))</td>
<td>UF Company, Costa Rica</td>
</tr>
<tr>
<td>EET 19, 95</td>
<td>Estación Experimental Tropical</td>
<td>10</td>
<td>The northeast of Ecuador (Coca River and north of Peru (Morona and Marañon rivers))</td>
<td>EET, Pichilingue, Ecuador</td>
</tr>
<tr>
<td>PA 150, 169</td>
<td>Parinari</td>
<td>16</td>
<td>Marañon River, Peru</td>
<td>ICG,T</td>
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<tr>
<td>SCA 9,10</td>
<td>Scavina</td>
<td>19</td>
<td>Ucayali River, Peru</td>
<td>ICG,T</td>
</tr>
<tr>
<td>NA 33</td>
<td>Nanay</td>
<td>19</td>
<td>Nanay River, Peru</td>
<td>ICG,T</td>
</tr>
<tr>
<td><em>CRU 100</em></td>
<td>Cocoa Research Unit</td>
<td>19</td>
<td>Nanay River, Peru</td>
<td>ICG,T</td>
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<td>Imperial Collection Selection</td>
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<td>Western Amazonia</td>
<td>ICG,T</td>
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<td>Trinidad Selected Amazons</td>
<td>20</td>
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<td>SCA 11, 12, 6</td>
<td>Scavina</td>
<td>20</td>
<td>Ucayali River, Peru</td>
<td>ICG,T</td>
</tr>
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</table>
The cacao cultivar CCN 51 was characterized by the unique haplotype H21. This haplotype is separated by one mutational step from H22 in the mononucleotide motif repeat of CaCrSSR2 (Figure 4). H22 was also found in the International Cacao Collection in Catie, including the Matina accession, which is also a common cultivar. Another unique and closely related haplotype is H23, which was observed along with haplotype 22 in Iquitos, Peru. H22 and H23 formed a separate cluster (lineage) in the haplotype network (Figure 4).

H10 was the most common haplotype (42%) in a set of Trinitario cultivars followed by H20 (35%) and other less frequent haplotypes H5, H19, and H22 (Table 4).

4. DISCUSSION

4.1. Distribution of Haplotype Diversity

High haplotype diversity was observed in western Amazonia (Figure 3). This region represents a hot spot of cacao diversity (Thomas et al., 2012; Zhang & Motilal, 2016), where cacao populations have a high haplotype diversity in the river systems of Marañon, Amazon (Iquitos), Nanay, and Uyacali in Peru, and along the Coca and Napo river systems in Ecuador (Figure 2 and Figure 3).

Our results support the hypothesis of decreasing cacao diversity from the western part to the eastern part of the range. The same differentiation pattern from west to east was observed in the Amazon basin for 200 cacao accessions representing ten genetic groups based on whole genome sequencing and multidimensional scaling analysis (Cornejo et al., 2018). Thus, the distribution of haplotype diversity is consistent with the suggested origin of the species in

<table>
<thead>
<tr>
<th>Clone name</th>
<th>Source</th>
<th>Haplotype</th>
<th>Tentative geographic origin</th>
<th>Breeding program origin</th>
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<tbody>
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<td>Imperial Collection Selection</td>
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<td>Western Amazonia</td>
<td>ICG.T</td>
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<tr>
<td>IMC 47, 60, 67</td>
<td>Iquitos Mixed Calabacillo</td>
<td></td>
<td>Iquitos, Peru</td>
<td>ICG.T</td>
</tr>
</tbody>
</table>

Accession name refers to an established accession name accepted at national and international level. * Trinitario cultivars; CATIE: Centro Agronómico Tropical de Investigación y Enseñanza, Costa Rica; ICGT: International Cocoa Genebank, Trinidad. Agronomic details (yield, disease resistance, and favorable traits) of these cacao clones are available in The International Cocoa Germplasm Database (ICGD) (Turnbull & Hadley, 2019).
western Amazonia (Bartley, 2005; Cheesman, 1944; Cuatrecasas, 1964; Motamayor et al., 2008).

Our results support the hypothesis of western Amazonia being considered as a forest refugial area (Haffer & Prance, 2001; Prance, 1982) and a center of crop genetic diversity (Clement, 1989; Clement et al., 2015), where cacao populations could have been restricted to temporarily isolated refugia during the Pleistocene (Figure 1). After the Pleistocene, suitable habitats during the Holocene allowed migration of cacao populations by human dispersal from western to central Amazonia along the Amazon River (Thomas et al., 2012). Cacao populations may have experienced selection in this center of crop diversity to some degree by domestication for fruit pulp and later were dispersed through the basin by pre-Columbian human expansion (Clement et al., 2010; Clement et al., 2015). Western Amazonia may also be a center of domestication which is reflected in the presence of predominant haplotypes H10, H20, and H22 both in breeding populations and plantations in western Amazonia (Figure 2 and Table 3).

Haplotypes that were observed only in western Amazonia (H12 in Colombia, H13 in Ecuador, and H11, H3, H2, H14, H15, H18, and H23 in Peru) probably are associated with cacao populations that experienced isolation in forest refugia when the Amazon forest was reduced during the Pleistocene (Figure 1) (Prance, 1982; Thomas et al., 2012; van der Hammen & Hooghiemstra, 2000). Another possible explanation assumes in situ occurrence due to edaphic adaptation to rich soils of refugial areas, which is common in tree species of Amazonia (Gentry, 1982, 1989).

Although the cacao samples studied here were distributed across the Amazon basin, they mainly represented the Upper Amazon (Figure 1). Thus, it is important to analyze additional cacao populations within the species range that could reveal new and distinctive haplotypes. For example, a recent forest survey of 1170 plots in Amazonia identified cacao as a common
tree in the basin; specifically, cacao populations were observed in southwestern and southern Amazonia, and French Guyana (Levis et al., 2017; Steege et al., 2013). Pre-Columbian human societies likely enriched these forest areas with cacao trees brought from the center of cacao diversity in western Amazonia (Levis et al., 2017).

The occurrence of only a single haplotype H16 in French Guiana could indicate human mediated dispersal of a few cacao individuals to this region. However, additional evidence to support human-mediated distribution of cacao in Amazonia is needed using more extensive sampling, specific studies such as radiocarbon and stable isotope analyses, and archaeological and ecological surveys looking at past human interactions with tropical tree species (Caetano-Andrade et al., 2020; Levis et al., 2018).

4.2. Chloroplast Haplotypes Match Genetic Groups Based on nSSRs

Haplotype diversity and distribution observed in our study partly mirrored the genetic clusters of the dendrogram based on 96 nSSRs (Figure 2 in [2]). For example, the Marañón, Guiana and Amelonado cluster is represented by related haplotypes 16 and 17 in the haplotype network (Table 4, Figure 4). Moreover, the cluster comprising Nanay and Iquitos is represented by the related haplotypes 19 and 22, and the closely related clusters Curaray and Nacional by the related haplotypes H10 and H9 in our haplotype network (Table 4, Figure 4).

The presence of specific haplotypes in different geographic regions and within different genetic groups suggest particular patterns of seed dispersal. For example, haplotype H20 was observed in Curaray (north of Ecuador), Iquitos (north of Peru), and Contamana (central Peru) groups (Figure 3, Table 4). Similarly, haplotype H19 was observed in the distantly related Contamana (central Peru) and Nanay (north of Peru) groups (Figure 3, Table 4). This may be explained by the presence of temporarily separated forest refugia in western Amazonia (Clement, 1989; Prance, 1982) where cacao populations were likely restricted during the
Pleistocene (Thomas et al., 2012). Later, during the Holocene cacao populations experienced habitat expansion when the climate became wetter and warmer, favorable for wide distribution in the Amazon basin (Thomas et al., 2012). Cacao habitat expansion aided by human mediated dispersal during the Holocene times may explain ample seed dispersal observed among cacao groups (Levis et al., 2017; Thomas et al., 2012).

Marañon, Guiana and Amelonado clustered together in the nSSR-based dendrogram (Motamayor et al., 2008); these populations share H16, and Amelonado and Marañon also have the related haplotype H17 in common (Table 4). These haplotypes reveal a wide distribution from western to eastern Amazonia (Figure 2) and likely experienced human dispersal in the basin. Indeed, current abundance of cacao populations in south-southwestern Amazon and French Guyana was explained partially by pre-Columbian human dispersal (Levis et al., 2017).

Cacao populations in French Guyana probably originated from local forest refugia during the Pleistocene–Holocene epochs (Lachenaud & Zhang, 2008). However, an alternative explanation of their origin may be human meditated seed dispersal from western to eastern Amazon (Levis et al., 2017; Sereno et al., 2006; Thomas et al., 2012). The observed wide distribution of haplotypes of H16 and H17 in the Amazon basin supports this hypothesis (Figure 2 and Figure 3). However, to clarify the origin of Amelonado and Guiana groups more extensive sampling across the species’ geographic range is needed accompanied by historical records of cacao dispersal in the Amazon. However, difficulties may arise to identify the hypothetical origin of native Amazonian crops due to forest expansion during the Holocene (Clement et al., 2010).

In the case of the Nacional group, two main natural populations with H10 were observed: one in the Ecuadorian Amazon (Napo River) and another in the Peruvian Amazon (Morona River) (Figure 2). We consider that additional cpDNA analyses in Ecuadorian populations are
needed to provide more evidence about the origin of the Nacional variety. However, our results are in agreement with (Motamayor et al., 2008) who found 13 individuals from the Morona River clustered in the Nacional group. Furthermore, based on the nSSR genetic analysis of 65 wild individuals widespread in the Amazonian forest of Ecuador and eight individuals from the Morona River in Peru, (Loor Solorzano et al., 2012) suggested an ancestral origin of the Nacional group in the Southern region of Ecuadorian Amazon close to Morona River.

The distribution of H10 suggests a pattern of seed dispersal among cacao populations in western Amazonia, which supports the hypothesis of a shared origin of this haplotype from Ecuadorian and Peruvian populations and later their introduction to the coast of Ecuador, where they were established in pure plantations. Furthermore, chloroplast analysis of Chuncho, a cacao population from Peru that resembles the Nacional fine-flavor cacao (Eskes et al., 2018), would help to trace the geographic origin of fine-flavor cacao to the south of Peru as suggested by (Eskes et al., 2018).

4.3. Chloroplast Haplotypes and Domesticated Cacao

Combined analyses of nuclear and cpDNA variation can help to unravel the history of cacao domestication and to narrow down the origin of domesticated varieties. For example, the Criollo variety (not included in the present study) was domesticated from a fraction of the ancestral Curaray population about 3600 years ago (Cornejo et al., 2018). Within the Curaray population we observed haplotypes H9, H13, and H20 in northern Ecuador (Figure 3 and Table 4). CpDNA analyses of the Criollo variety could reveal its geographic origin within the distribution range of Curaray. Indeed, the geographic origin of haplotype H9 observed with high frequency in the Curaray group (Figure 4) possibly suggests that trees from the Ecuador-Colombia border served as a source of plant material during the domestication of Criollo cacao (Figure 3).
Further questions arise such as why the domestication of cacao started with the Ecuadorian Curaray group and not with other populations. It was probably because its sweet pulp was used for the elaboration of fermented beverages, which may also have favored its selection and dispersal out of the Ecuadorian Amazon (Clement et al., 2010; Loor Solorzano et al., 2012; Zarrillo et al., 2018).

4.4. Haplotype Diversity in Plantations and Breeding Populations

The high haplotype diversity observed in natural populations contrasts with the low diversity observed in cacao plantations (Table 2). We observed a high contribution of haplotype H10 in both selections made in cacao plantations in the west of Ecuador and breeding populations such as the ICS clones of Trinidad (Table 3). In addition, H10 was also observed in Colombia and Grenada, providing further evidence for introduction events of this haplotype to other countries.

Cacao cultivation in coastal Ecuador has been recorded since the 17th century, and cacao genotypes named Nacional were selected and cultivated because of their special chocolate flavor (Bartley, 2005). Currently, new sources of genetic material are required to improve flavor, increase yield and add disease resistance traits to the Nacional cultivar (Loor Solorzano et al., 2012). We suggest that the four wild provenances associated with haplotype H10 in western Amazonia of Peru and Ecuador could be screened for agronomic traits to test their use in breeding programs, potentially adding useful traits to Nacional cultivars.

The presence of six haplotypes, with haplotype H10 being the most frequent (42%), in 15 reference Trinitario accessions suggests seed introduction events occurring probably from northeast of Ecuador and north of Peru to Trinidad (Table 3). Similar to our results, five different haplotypes were observed in 21 different reference Trinitario accessions based on nine cacao cpSSRs markers (Yang et al., 2013).
The river basins associated with geographically restricted haplotypes (Figure 2 and Figure 3) could serve as source of specific adaptations and potential new traits for cacao breeding programs, but only if these samples carry favorable agronomic traits. Examples of these areas include the Napo basin of Ecuador (H5, H9, and H13), the Caquetá River (H12) in Colombia, and Ucayali River in south and central Peru (H1).

5. CONCLUSIONS

Western Amazonia contains high haplotype diversity areas in Peru and Ecuador. Populations with at least two common haplotypes may have been dispersed by humans from this center of cacao genetic diversity to new suitable habitats following a west to east route of migration in the Amazon basin. The western Amazonia has high value for cacao conservation considering its haplotype diversity and presence of geographically unique haplotypes, which were not observed in breeding populations and cultivars yet and could be evaluated for agronomic traits in support of cacao improvement.

The cacao cpDNA haplotypes observed here can be used to determine the chloroplast diversity of accessions and select distinctive haplotypes in cacao breeding programs. Additionally, the map of cpDNA haplotypes can be used to verify the geographic origin of planting material at finer geographic scale and point to potential areas for new collections. Craft chocolate makers can use this reference map to create new markets for cacao products based on geographic origin.

6. OUTLOOK

New collections in Amazonia will expand the availability of cacao genetic resources and help us better understand the patterns of geographic genetic variation and dispersal. However, rapid land use changes and gold mining in the Amazon forest are threatening the natural habitat of wild cacao populations and eroding its genetic diversity (Sierra, 2000; Steege et al., 2013;
Zhang et al., 2011). Likewise, examples of documented cacao natural populations that may reveal new haplotypes are Upper Orinoco (Patiño, 1963), Caquetá River in Colombian Amazon and Ecuadorian Amazon (Allen, 1988), Santiago and Morona rivers in northern Peru (Arevalo-Gardini et al., 2019), Chuncho in southern Peru (Eskes et al., 2018), Beni River in Bolivia (Zhang et al., 2012), the Brazilian Amazon (Almeida et al., 2015; L.A.S Dias et al., 2003; Sereno et al., 2006), southwestern and southern Amazonia (Levis et al., 2017; Wittmann et al., 2006), and Amazonian basin (Motamayor et al., 2008). Furthermore, the analysis of additional samples like those held in international (CATIE and ICGT) and national cacao collections (see Turnbull & Hadley, 2019) likely will reveal additional chloroplast diversity.

AUTHOR CONTRIBUTIONS


FUNDING

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DATA AVAILABILITY STATEMENT

The analyzed germplasm are available on request to The International Cocoa Quarantine Centre at the University of Reading (ICQC,R) (Turnbull & Hadley, 2019).
ACKNOWLEDGMENTS

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CONFLICTS OF INTEREST

The authors declare no conflict of interest.
7 REFERENCES


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8 SUPPLEMENTARY MATERIALS

The following are available online at https://www.mdpi.com/article/10.3390/d13060249/s1.

**Table S1**: Clone name, ICQC (The International Cocoa Quarantine Centre) accession number, geographic origin, samples per location, genetic group according to Motamayor et al. 2008, and chloroplasts haplotypes observed in a sample of *Theobroma cacao* L. accessions. The haplotype column shows the order of the markers *CaCr1, CaCr2, CaCr4, CaCr5, CaCr8* and *CaCr9*, and the fragment sizes are given in base pairs (bp). Details about donor collections, geographic location of collections expeditions and agronomic traits of these cacao clones are available on the ICGD (The International Cocoa Germplasm Database) website (Turnbull & Hadley, 2019).

**Table S2**: Clone name, ICQC accession number, locations, and chloroplast haplotypes of *Theobroma cacao* L. in nine farms of the pacific coast of Ecuador. The haplotype column shows the order of the markers *CaCr1, CaCr2, CaCr4, CaCr5, CaCr8* and *CaCr9*, and the fragment sizes are given in base pairs (bp). Collection and agronomic details of these cacao clones are available on the ICGD website.

**Table S3**: Clones names, ICQC accession number, origin of samples, genetic group (Motamayor et al. 2008), and chloroplasts haplotypes observed in a sample of *Theobroma cacao* L. genotypes used for cultivation or breeding. The haplotype column shows the order of the markers *CaCr1, CaCr2, CaCr4, CaCr5, CaCr8* and *CaCr9*, and the fragment sizes are given in base pairs (bp). Agronomic details of these cacao clones (yield, disease resistance, and favorable traits) and breeding details are available on the ICGD website.
Paper 3

Genotyping of cacao (*Theobroma cacao* L.) germplasm resources with SNP markers linked to agronomic traits reveals signs of selection

Submitted

*Tree genetics & Genomes*
Genotyping of cacao (*Theobroma cacao* L.) germplasm resources with SNP markers linked to agronomic traits reveals signs of selection

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**Abstract**

The Amazonian cacao tree or the chocolate tree (*Theobroma cacao* L.) is cultivated to produce seeds, the valuable raw material for the chocolate industry. However, cacao yield is hampered by diseases and low productivity. Cacao SNP markers associated with breeding and adaptive traits provide a genetic tool for improving selection and reducing breeding cycles. Here, we genotyped 40 published SNPs associated with disease resistance and 11 SNPs with yield traits in 346 accessions using the MassARRAY® system. Four genetic clusters were identified, and two of them were observed in high proportion in managed cacao, characterized by a high proportion of admixed individuals reflecting the man-made hybrids. One cluster overrepresented in managed cacao is associated with the preference to keep superior flavor in cultivated cacao from Criollo ancestry. Introgression of wild material collected in Peru associated with disease resistance to witches’ broom disease (WBD) was also observed.
Underutilized genetic resources were observed in managed cacao such as the Guiana cacao, while previous agronomic evaluation has demonstrated its good yield potential. Yield and disease resistance traits (mainly resistance to WBD) show divergence between wild and managed cacao ($F_{ST} > 0.05$) probably reflecting selection during domestication, cultivation and breeding efforts. The identified SNPs showing divergence between wild and managed cacao can be used to build breeding populations. New collections of wild cacao followed by agronomic evaluations can broaden the genetic base of cultivated cacao, especially for such traits as disease resistance to frosty pod rot (FPR).

**Keywords:** Amazonia, cacao, disease resistance, germplasm resources, SNPs, tropical crop breeding.
1 INTRODUCTION

The cacao (Theobroma cacao L.) is a valuable crop tree commonly cultivated in the tropics to produce seeds, the raw material that sustains the global chocolate industry (Coe and Coe 2013; Dand 2011). The putative origin of cacao is western Amazonia (Upper Amazon basin), and its natural range includes the wide Amazon basin, where the primary gene pool of cacao is structured in twelve genetic clusters (Gutiérrez et al. 2021a; Motamayor et al. 2008; Nieves-Orduña et al. 2021; Nieves-Orduña et al. 2023; Sereno et al. 2006; Zhang et al. 2011, 2012). Seven of these genetic clusters were identified in natural populations in western Amazonia, at the Ecuadorian-Peruvian Amazon, whereas one cluster is dominant in eastern Amazonia and French Guiana (Cornejo et al. 2018; Motamayor et al. 2008). Further population genetic analysis on areas not yet studied can identify new clusters and potential genetic resources for breeding (Nieves-Orduña et al. 2023). Cacao populations from western Amazonia were likely spread by humans to Eastern Amazonia (Levis et al. 2017). This dispersal generated a gradient of cacao genetic diversity declining from west to east Amazonia (Cornejo et al. 2018; Motamayor et al. 2008; Nieves-Orduña et al. 2021). In addition, the influences of paleoclimates and forest refugial areas also shaped the distribution of cacao in Amazonia (Lachenaud 1997; Motamayor et al. 2008; Thomas et al. 2012).

The cacao genetic resources are conserved mainly in two international and several national collections (Bekele and Phillips-Mora 2019). Continued deforestation in wild habitats threatens wild cacao diversity if proper in situ conservation and new collections to enrich current ex situ collections are not implemented (Nieves-Orduña et al. 2023). The International Cocoa Quarantine Centre at the University of Reading (ICQC, R) in the UK facilitates the global distribution of pathogen-free plant material (Daymond 2018), but cultivated cacao represents only a fraction of the species’ genetic diversity (Bennett 2003; Boza et al. 2014; Zhang et al. 2011; Zhang and Motilal 2016). The global demand for
chocolate has increased cacao cultivation, but it is hampered by diseases and low yield (Gutiérrez et al. 2016; Ploetz 2016). In addition, there is evidence of deforestation linked to cacao expansion mainly in west Africa (Hoang and Kanemoto 2021; Kalischek et al. 2023) but also increased forest cover through agroforestry (Orozco-Aguilar et al. 2021). Compared to advances in the productivity of tropical crops, such as oil palm, average global cacao production has remained low since 1961 (Morrissey et al. 2019).

Advances in cacao molecular marker assisted breeding and genomic selection can improve breeding populations and help to select high yield and disease-resistant cacao genotypes (Schnell et al. 2007). Still cacao breeding depends on the extensive genetic variation existing in wild populations present in Amazonia (Motamayor et al. 2008; Nieves-Orduna et al. 2023). Genome-wide association studies (GWAS) and quantitative trait locus (QTL) mapping in cacao have identified single nucleotide polymorphism (SNP) markers linked to resistance to the black pod rot (BPR) disease caused by *Phytophthora* spp. (Gutiérrez et al. 2021b), witches’ broom disease (WBD) caused by *Moniliophthora perniciosa* (Motilal et al. 2016; Royaert et al. 2016), Ceratocystis wilt (CW) caused by *Ceratocystis cacaofunesta* (Fernandes et al. 2018), and moniliasis or frosty pod rot (FPR) caused by *Moniliophthora roreri* (Gutiérrez et al. 2021b). In addition, SNPs linked to high yield traits and sexual compatibility were identified (da Silva et al. 2016; Fernandes et al. 2020). Thus, favorable identified genetic variation can be exploited for improving the selection efficiency and reducing breeding cycles.

We present here our research aimed to characterize a diverse set of 346 reference cacao accessions representing both wild and managed cacao genotyped for 51 published SNPs associated with important cacao agronomic traits, such as disease resistance, yield and sexual compatibility, using the MassARRAY® system (Agena Bioscience, Hamburg, Germany). Specifically, our aims were to 1) describe the population structure and analyze genetic clusters in wild and managed cacao, 2) identify new genetic resources for cacao breeding based on SNP
profiles, and 3) identify signatures of selection for agronomic traits that differentiate wild and managed cacao to assist breeding of superior cacao genotypes. Using published phenotypic data, we aimed to also validate disease resistance and yield associated SNP alleles and uncover valuable cacao accessions and new genetic resources to be used in breeding efforts. The obtained SNP profiles will be reported to the International Cocoa Germplasm Database (ICGD). The accessions analyzed here are in the public domain and can be accessed by any cacao-producing country, facilitating the validation of results through breeding programs and farm conditions.

2 MATERIALS AND METHODS

Germplasm and DNA Isolation

The germplasm analyzed consisted of 346 cacao accessions collected in wild habitats and from managed cacao. The managed cacao included 168 clones representing different cultivars, selections and breeding populations from several groups such as Refractario, Estación Experimental Tropical (EET), United Fruit (UF), Imperial College Selection (ICS), Trinidad Select Hybrid (TSH), SIAL (Selecao Instituto Agronomico do Leste), Selecao Instituto do Cacau (SIC) and Tropical Agriculture Research Service (TARS-Series of cacao) (Turnbull and Hadley 2023). The wild germplasm samples included 178 accessions, representing mainly northwestern Amazonia (77% of the samples), which is considered the hot spot of cacao genetic diversity (Clement et al. 2010; Cornejo et al. 2018), and French Guiana (23%). The wild cacao germplasm represented a wide geographic distribution within the cacao primary gene pool, including samples from five countries, mostly from Peru (57%). They included accessions known as the Pound Collection composed of groups coded as IMC, MO, NA, PA, POUND, and SCA. These accessions were collected in the 1930s by Frederick J. Pound in the Peruvian Amazon while searching for genotypes resistant to WBD that are now the basis for breeding WBD-resistant cacao (Bartley 2005; Zhang et al. 2011). More recent collections included cacao
accessions coded as LCT EEN collected in the Ecuadorian Amazon (Allen 1988) and GU accessions from French Guiana (Lachenaud 2015). Although collected in the wild as seeds or budwood, these groups of accessions may have experienced human intervention such as cultivation or translocation, considering the persistent effects of pre-Columbian societies in plant domestication in Amazonia (Barlow et al. 2012; Clement et al. 2015; Levis et al. 2017). Among the accessions analyzed, 137 clones represented the ten genetic groups identified by Motamayor et al. (2018): Criollo (3), Amelonado (22), Nacional which represent traditional cultivars (2), Curaray (10), Contamana (5), Guiana (15), Iquitos (30), Marañon (32), Nanay (10) and Purus (8). The complete list of samples studied is presented in Supplementary Table S1. In addition, the ICGD (http://www.icgd.rdg.ac.uk) provides updated agronomic details, geographic origin, and passport data of the germplasm analyzed (Turnbull and Hadley 2023).

The 346 cacao accessions analyzed are subject to international distribution under the International Treaty on Plant Genetic Resources for Food and Agriculture (CacaoNet 2012) and were obtained thanks to the ICQC at the University of Reading, UK., and the germplasm cacao collection held in the International Center for Tropical Agriculture (CATIE) in Costa Rica. We used 1 cm$^2$ of fresh leaf tissue per sample and the DNeasy 96 Plant Kit (Qiagen, Hilden, Germany) for DNA extraction.

**Selection of the SNP markers**

We selected 51 cacao SNPs from the published data that are supposedly associated with important agronomic traits such as disease resistance (40 SNPs) and yield (11 SNPs) and genotyped them in wild and managed cacao accessions in our study (Table 1). This set of SNPs was identified by GWAS and QTL analysis in cacao research centers at Trinidad (Motilal et al. 2016), Costa Rica (Gutiérrez et al. 2021b) and Brazil (da Silva et al. 2016; Fernandes et al. 2018; Fernandes et al. 2020; Royaert et al. 2016). The SNP panel included eight SNPs associated with disease resistance to BPR (Gutiérrez et al. 2021b), 11 SNPs linked to WBD
resistance (Motilal et al. 2016; Royaert et al. 2016), 16 SNPs related to FPR resistance (Gutiérrez et al. 2021b), and five SNPs linked to CWC resistance (Fernandes et al. 2018). In addition, the SNP set included three SNPs associated with number of seeds (Motilal et al. 2016), four SNPs related to yield components such as dry seed weight, number of pods harvested, average yield and high pod index, respectively (Fernandes et al. 2020), and four SNPs associated with flower retention (as a measure of self-compatibility) (da Silva et al. 2016).

Table 1 Fifty-one SNPs associated with agronomic traits (disease resistance and yield) in *Theobroma cacao* L. and genotyped in the present study in 346 cacao accessions

<table>
<thead>
<tr>
<th>Agronomic trait (reference)</th>
<th>SNP ID</th>
<th>Identification method and study population</th>
</tr>
</thead>
<tbody>
<tr>
<td>Witches’ broom disease, WBD (Royaert et al. 2016)</td>
<td>Tcm003s33466269, Tcm004s00110232, Tcm006s19715703, Tcm006s25375496, Tcm007s10302466, Tcm009s08066239, Tcm009s02031341</td>
<td>QTL mapping: 1) F₁ population (459 trees) at Mars Center for Cocoa science, Bahia, Brazil; 2) TSH 1188 (resistant) × CCN 51 (tolerant)</td>
</tr>
<tr>
<td>Witches’ broom disease, WBD (Motilal et al. 2016)</td>
<td>TcSNP720, TcSNP1230, TcSNP375, TcSNP1374</td>
<td>Genome wide associating mapping: 483 unique accessions from the International Cocoa Genebank Trinidad (ICGT)</td>
</tr>
<tr>
<td>Ceratocystis wilt, CW (Fernandes et al. 2018)</td>
<td>Tcm006s13222057, Tcm006s13371871, Tcm006s1372133, Tcm004s02243097, Tcm004s0274866</td>
<td>QTL mapping: 1) F₁ population (266 trees) at Mars Center for Cocoa science, Bahia, Brazil; 2) TSH 1188 (resistance) × CCN 51 (susceptible)</td>
</tr>
<tr>
<td>Frosty pod rot (FPR) or Moniliasis (Gutiérrez et al. 2021)</td>
<td>Tcm004s01757744, Tcm004s01737820, Tcm007s05825365, Tcm007s04093978, Tcm008s05308021, Tcm008s05903121, Tcm009s04190398, Tcm009s04191545, Tcm010s02288537, Tcm010s02455243, Tcm002s04088162, Tcm002s04156143, Tcm009s04033054, Tcm009s040465466, Tcm010s00936105, Tcm010s00913858</td>
<td>QTL mapping: 1) F₁ population (179 trees) at CATIE, Costa Rica; 2) POUND 7 (moderately susceptible to FPR and resistant to BPR) × UF 273 (resistant to FPR and highly susceptible to BPR)</td>
</tr>
<tr>
<td>Black pod rot, BPR (Gutiérrez et al. 2021)</td>
<td>Tcm002s08313597, Tcm002s08269115, Tcm004s28538741, Tcm004s26962336, Tcm008s04656460, Tcm008s04576575, Tcm010s22418501, Tcm010s21842721</td>
<td></td>
</tr>
<tr>
<td>Dry bean weight/yield, number of pods harvested (Fernandes et al. 2020)</td>
<td>Tcm004s00289192, Tcm004s00615809</td>
<td></td>
</tr>
<tr>
<td>Average yield (Fernandes et al. 2020)</td>
<td>Tcm004s01127580</td>
<td></td>
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<tr>
<td>Pod index (Fernandes et al. 2020)</td>
<td>Tcm002s23708704</td>
<td></td>
</tr>
</tbody>
</table>
Agronomic trait (reference) | SNP ID | Identification method and study population
--- | --- | ---
Number of seeds (Motilal et al. 2016) | TcSNP1370, TcSNP697, TcSNP368 | Genome wide associating mapping: 483 unique accessions from the International Cocoa Gene bank Trinidad (ICGT)

Flower setting, as a measure of self-compatibility (da Silva et al. 2016) | TcSNP1866, TcSNP1867, TcSNP1869, TcSNP1870 | Genome wide associating mapping: 1) 295 unique accessions from the germplasm collections at Cacao Research Center (CEPEC), Bahia, Brazil; 2) 256 randomly selected individuals from a third breeding generation.

**Genotyping with MassARRAY system**

The SNPs’ flanking sequences were obtained from the published data (see Table 1) and were used to design two SNPs assays for the MassARRAY® system using the Assay Design Suite V2.0 (Agena Bioscience 2015). Primer adjustment, PCR amplification, SAP treatment and iPLEX reaction were done following the instructions of the manufacturer (Agena Bioscience 2019a). Allele calling was conducted using Typer Analyzer v.5.0.2137 (Agena Bioscience 2019b). SNPs with at least 75% genotyping success call rate across all samples were retained for further analysis, the same genotyping success call rate threshold was used for all DNA samples. Thus, the final data set for further analysis included 318 DNA samples and 42 SNPs (Supplementary Tables S1, S2 and S3).

**Data analysis**

Analysis of molecular variance (AMOVA) and $F_{ST}$ between wild and managed cacao, $F_{ST}$ per locus, and principal coordinate analysis (PCoA) were performed in GenAlEx 6.5 using 999 permutations (Peakall and Smouse 2012). Analysis of population structure and admixture was done using STRUCTURE 2.3.4 (Pritchard et al. 2000) with the admixture model and correlated allele frequencies of 42 SNPs by testing from $K = 1$ to $K = 10$ subpopulations with 10 repetitions for each $K$. The numbers of burn-ins and iterations were 10,000 and 100,000, respectively. Structure Harvester 0.6.94 (Earl and vonHoldt 2012) was used to determine the
most likely number of clusters (K) using the K method, and obtained results were visualized using CLUMPAK (Kopelman et al. 2015).

An UPGMA dendrogram based on the pairwise Nei’s standard genetic distance matrix of individuals genotyped for 42 SNPs was generated using Populations 1.2.32 (Langella 2001) with 999 bootstraps on loci and visualized using Interactive Tree of Life (iTOL) 6.7.4 (Letunic and Bork 2021).

Phenotypic trait data to validate favorable SNPs associated with agronomic traits were collected from published data and the International Cocoa Germplasm Data Base (Turnbull and Hadley 2023). Based on these published data, accessions were described as either susceptible or tolerant/resistant, while no clear distinction was made between the terms tolerant and resistant. Fisher’s exact test was calculated in Statistica (StatSoft Europe GmbH, Hamburg, Germany) to compare genotype distributions of wild and managed cacao, and of resistant-tolerant and susceptible germplasm to WBD and CW. Relative genotype frequencies per SNP were calculated in Excel (Microsoft Corporation, Redmond, Washington, USA). The map showing the geographic distribution of identified cacao clusters was developed using ArcGIS software (www.esri.com).

3 RESULTS

Genetic structure of accessions observed in wild and managed cacao

Although the structure analysis did not resolve the exact ten genetic clusters identified earlier in cacao populations based on microsatellite markers (Motamayor et al. 2008), it suggests four most likely clusters (K = 4) for the studied accessions based on 42 SNPs (Figure 1 and Supplementary Figures S1-S2).

We observed a clear pattern of differentiation of the wild germplasm when it was arranged from west to east Amazonia. Accessions collected in Ecuadorian Amazon showed a high admixture proportion of cluster two (Figure 1A). Cluster one was observed in high proportion
in the Peruvian Amazon. The IMC, NA, and PA germplasm series collected around Iquitos, Peru have similar structure and are dominated by cluster one, but the PA series also admixed much with the fourth cluster (Figure 1B). The MO series collected in Peru have a high admixture from the third cluster, which has a high representation in managed cacao. The SCA series showed a high proportion of the second cluster. The accessions collected in French Guiana can be distinguished by a high admixture from the fourth cluster (Figure 1). No private alleles were observed in the data set.

We observed that managed cacao accessions have mainly admixture from the second and third clusters, with a little admixture from the first cluster (Figure 1). A high proportion of the

Fig. 1 Genetic admixture of four clusters identified in wild and managed cacao accessions based on 42 SNPs. a - geographic distribution of clusters 1-4 in wild cacao. b – Q-values of individual wild cacao accessions arranged along their geographical location from west to east Amazonia. c - Q-values of individual managed cacao accessions arranged from high to low admixture of the second cluster. MO (Morona), IMC (Iquitos Mixed Calabacillo), NA (Nanay), POUND, PA (Parinari) and SCA (Scavina) are the names of groups of accessions from the Pound Collection.
second cluster was observed in the Criollo accessions (Criollo 12, 13, 65), which represent the first domesticated cacao (Cornejo et al. 2018). A higher proportion of the third cluster was observed in clones developed in Brazil, such as SIAL and SIC. In addition, we identified underrepresentation of the fourth cluster associated with Guiana accessions. Overall, managed cacao includes largely admixed accessions, reflecting hybrids in breeding populations and cultivars such as UF, ICS, TARS, CCN 51 and CATIE R6 (Figure 1 and Supplementary Figure 2B).

Fig. 2 Principal coordinates analysis (PCoA) of 318 cacao accessions based on genetic distance matrix estimated with 42 SNP markers linked to agronomic traits. The first component of the PCoA explains 13.09% of the total variation, and the second component explains 8.31%. Different dot colors represent different wild accessions; open green circles depict managed cacao accessions.

Although there is an overlap between wild and managed cacao in the PCoA, we observed some clustering trends in wild cacao and managed cacao (Figure 2). Geographically, accessions collected in eastern Amazonia, GU-Guiana, are clustered in the lower left of the PCoA. While
wild accessions collected in western Amazonia tend to be clustered on the right side of the PCoA. Accessions such as SCA collected in central Peru are located in the lower right of the PCoA (Figure 2). The dendrogram shows a similar pattern as the PCoA, the Guiana accessions are separated as a separate group, sister to another group represented by a subset of accessions of managed cacao. The rest of the accessions of managed cacao is not well-resolved in the dendrogram (Figure 3).

**Fig. 3** UPGMA dendrogram of 318 cacao accessions based on the Nei’s standard genetic distance of individuals estimated with 42 SNPs linked to agronomic traits. Clades supported by significant bootstrap values (above 50%) are shown with respective values.

Managed cacao accessions tend to be clustered in the upper left of the PCoA, specially selected cacao such as SIC and SIAL developed in Bahia Brazil, but Criollo accessions (Criollo 12, 13, 65) are clustered on the right side. Within the managed cacao, cultivars and selections of high use in breeding programs such as CCN 51 and UF 273 type 1 are located in the center.
of the PCoA. CCN 51 is plotted closely to RB 39, which is resistant to WBD. Wild accessions genetically close to UF 273 type 1 include PA series, such as PA 169 and PA 4. PA 4 has resistance to WBD and BPR, and PA 169 is commonly used as a source of resistance to BPR and FPR (Turnbull and Hadley 2023). The accessions IMC 6 and IMC 31 are both characterized for having a low pod index (high yield) and resistance to Phytophthora (Turnbull and Hadley 2023) and are closely plotted to UF 273 type 1 in the PCoA (Figure 2).

Agronomic traits showing divergence between wild and managed cacao

AMOVA analysis indicated that 5% of the total variation was caused by variation between wild and managed cacao, 37% by the variation among individuals and 58% by the variation within individuals (Table 2). In addition, 11 SNPs showed $F_{ST}$ values above 0.05, indicating moderate genetic differentiation between managed and wild cacao. Six of these SNPs had $F_{ST}$ values greater than 0.10, suggesting strong divergence between the two groups at these markers (Table 3).

Table 2 Analysis of molecular variance (AMOVA) in the genotyped managed and wild cacao groups

<table>
<thead>
<tr>
<th>Source</th>
<th>d.f.</th>
<th>Sum of squares</th>
<th>Means squares</th>
<th>Est. var.</th>
<th>% of total variation</th>
<th>$F_{ST}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Among groups</td>
<td>1</td>
<td>144,295</td>
<td>144,295</td>
<td>0.415</td>
<td>5</td>
<td>0.046*</td>
</tr>
<tr>
<td>Among individuals</td>
<td>316</td>
<td>3886.551</td>
<td>12.299</td>
<td>3.470</td>
<td>37</td>
<td></td>
</tr>
<tr>
<td>Within individuals</td>
<td>318</td>
<td>1704.000</td>
<td>5.358</td>
<td>5.358</td>
<td>58</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>635</td>
<td>5734.846</td>
<td>9.244</td>
<td>100%</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*P < 0.001

Table 3 Eleven SNPs associated with agronomic traits showing significant divergence between managed and wild cacao ($F_{ST} > 0.05$)

<table>
<thead>
<tr>
<th>Agronomic trait</th>
<th>SNP ID</th>
<th>$F_{ST}$</th>
<th>Chr.</th>
<th>Gene</th>
<th>Description</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flower setting (as a measure of self-compatibility)</td>
<td>TcSNP1866</td>
<td>0.260</td>
<td>4</td>
<td>n/a</td>
<td>n/a</td>
<td>n/a</td>
</tr>
<tr>
<td>Dry bean weight / high yield</td>
<td>Tc004s00289192</td>
<td>0.160</td>
<td>4</td>
<td>Thecc.04G002600</td>
<td>Family of uncharacterized protein function</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tc003s33466269</td>
<td>0.064</td>
<td>3</td>
<td>Thecc.03G307700</td>
<td>RNA-binding family protein</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tc004s00110232</td>
<td>0.101</td>
<td>4</td>
<td>Thecc.04G000500</td>
<td>UDP-glucose pyrophosphorylase 2</td>
<td></td>
</tr>
<tr>
<td>Witches’ broom disease (WBD)</td>
<td>Tc006s19715703</td>
<td>0.147</td>
<td>6</td>
<td>Thecc.06G104900</td>
<td>Hydroxyproline-rich glycoprotein family protein</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TcSNP1230</td>
<td>0.082</td>
<td>8</td>
<td>Thecc.08G127800</td>
<td>PDI- Protein disulfide isomerase1-1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Tc009s08066239</td>
<td>0.065</td>
<td>9</td>
<td>Thecc.09G141300</td>
<td>RING/U-box superfamily protein</td>
<td></td>
</tr>
<tr>
<td>Agronomic trait (CW)</td>
<td>SNP ID</td>
<td>$F_{ST}$</td>
<td>Chr.</td>
<td>Gene$^2$</td>
<td>Description$^2$</td>
<td></td>
</tr>
<tr>
<td>---------------------</td>
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<td>----------</td>
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<td></td>
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<tr>
<td>Ceratocystis wilt</td>
<td>Tcm006s13222057</td>
<td>0.123</td>
<td>6</td>
<td>Thecc.06G070400</td>
<td>Peptide deformylase 1A</td>
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<td>Pod index$^1$</td>
<td>Tcm002s23708704</td>
<td>0.107</td>
<td>2</td>
<td>Thecc.02G221200</td>
<td>Elongator protein 2</td>
<td></td>
</tr>
<tr>
<td>Moniliasis (FPR)</td>
<td>Tcm009s40465466</td>
<td>0.061</td>
<td>9</td>
<td>Thecc.09G335000</td>
<td>Pseudo-response regulator 7</td>
<td></td>
</tr>
<tr>
<td>Number of seeds</td>
<td>TcSNP1370</td>
<td>0.054</td>
<td>1</td>
<td>Thecc.01G096000</td>
<td>Histone superfamily protein</td>
<td></td>
</tr>
</tbody>
</table>

Note. All $F_{ST}$ values were significant at $P < 0.001$. Chr - chromosome. $^1$ Pod index: number of fruits needed to produce 1 kg of dry seeds; a low pod index (e.g., 14-20) is preferred because it helps to select trees that yield heavier and larger seeds (Bekele et al. 2020). $^2$ Gene ID and description based on Matina reference cacao genome (Theobroma cacao v2.1) (Motamayor et al. 2013) available at https://phytozome-next.jgi.doe.gov/info/Tcacao_v2_1

Three SNPs associated with yield traits showed high divergence between wild and cultivated cacao. Among these SNPs, Tcm004s00289192 ($F_{ST} = 0.160$, $P < 0.001$) was linked to dry bean weight and high yield, and TcSNP1370 ($F_{ST} = 0.054$, $P < 0.001$) was associated with number of seeds per fruit. Tcm002s23708704 ($F_{ST} = 0.107$, $P < 0.001$) was related to pod index, a measure of yield in commercial plantations. TcSNP1866 was associated with flower retention and had the highest $F_{ST} = 0.260$ ($P < 0.001$) between wild and managed cacao. Figure 4 shows the relative genotype frequency of SNPs showing pronounced divergence between wild and managed cacao and results of Fisher's exact test.

Managed and wild cacao showed also high divergence in disease resistance traits, especially in WBD resistance. We observed five SNPs significantly differentiated between managed and wild cacao that are associated with WBD resistance: Tcm006s19715703 ($F_{ST} = 0.146$, $P < 0.001$), Tcm004s00110232 ($F_{ST} = 0.101$, $P < 0.001$), TcSNP1230 ($F_{ST} = 0.083$, $P < 0.001$), Tcm003s33466269 ($F_{ST} = 0.067$, $P < 0.001$), and Tcm009s08066239 ($F_{ST} = 0.063$; $P < 0.001$). In addition, Tcm006s13222057 ($F_{ST} = 0.155$; $P < 0.001$) associated with resistance to CWC, and Tcm009s40465466 ($F_{ST} = 0.066$; $P < 0.001$) related to FPR resistance, were also significantly differentiated between managed and wild cacao (Table 3).
Fig. 4 Distribution of cacao genotypes for SNPs related to agronomic traits in the wild and managed cacao accessions. **a** - SNP markers related to yield traits: flower retention (da Silva et al. 2016); dry bean weight/high yield and high pod index (Fernandes et al. 2020). **b** - SNP markers related to disease resistance traits: Witches’ broom disease (Royaert et al. 2016) and Ceratocystis wilt (Fernandes et al. 2018). *Significant difference between wild and managed cacao based on the Fisher’s exact test ($P < 0.05$)

Figures 5 and 6 show the genotype frequencies for the SNPs Tcm004s00110232 and Tcm006s13222057 among cacao accessions resistant, tolerant and susceptible to WBD and CW, respectively. Germplasm with resistance (n=58) and tolerance (n=25) to WBD showed a significantly higher frequency of the CT and TT genotypes (Tcm004s00110232), and the CC genotype showed a low frequency in resistant plants, while all susceptible plants (n=9) showed the TT genotype (Figure 5). In addition, germplasm evaluated as resistant (n=28) and tolerant (n=16) to CW showed a higher frequency of GG and GT (Tcm006s13222057) genotypes, while the TT genotype was common in susceptible (n=21) plants. However, these differences were not statistically significant ($P > 0.05$, Figure 6).
**Fig. 5** Genotype frequencies for the SNP Tcm004s00110232 related to witches’ broom disease (WBD) among cacao (*Theobroma cacao* L.) accessions with resistance, tolerance, or susceptibility to WBD. Number of accessions is provided in brackets. Reaction to WBD reported by the International Germplasm Cocoa Database ([https://www.icgd.reading.ac.uk/index.php](https://www.icgd.reading.ac.uk/index.php)). *Significant difference between resistant-tolerant and susceptible based on the Fisher’s exact test (P = 0.0026)*

**Fig. 6** Genotype frequencies for the SNP Tcm006s13222057 associated with Ceratosystis wild disease among cacao (*Theobroma cacao* L.) accessions with resistance, tolerance, or susceptibility to Ceratosystis disease. Number of accessions is provided in brackets. Reaction to Ceratosystis disease reported by the International Germplasm Cocoa Database ([https://www.icgd.reading.ac.uk/index.php](https://www.icgd.reading.ac.uk/index.php)). No significant difference between resistant-tolerant and susceptible based on the Fisher’s exact test (P = 0.7910)
4 DISCUSSION

Low population structure in managed cacao

With our set of SNPs, we did not identify the ten genetic clusters observed earlier by Motamayor et al. (2008) based on 96 simple sequence repeat (SSR) markers. It is not surprising, considering a limited set of nonrandom 42 SNPs that we used in our study, which could be also under selection. However, we still identified four clusters reflecting the geographic origin of wild samples, with the second and third clusters being overrepresented in managed cacao, mainly consisting of hybrids (Figure 1). This population pattern in managed cacao is in agreement with observations made by Cornejo et al. (2018). They analyzed the genome sequence of 200 cacao accessions and identified that cultivated cacao and man-made hybrids are mainly composed of two clusters, Criollo and Amelonado, with a low admixture of the Nacional cluster (Cornejo et al. 2018). Population genetic analysis demonstrated that Criollo was the first domesticated cacao and provides the foundations of cultivated cacao until today, mostly due to flavor and chocolate attributes (Lachenaud and Motamayor 2017; Motamayor et al. 2002). Our results confirm that a narrow cacao genetic diversity has been used in managed cacao likely due to retaining quality traits derived from Criollo observed in the second cluster in our analysis.

Cornejo et al. (2018) also observed that a higher Criollo ancestry in man-made hybrids is associated with low yield (seed productivity per year per plant) mainly due to the accumulation of deleterious mutations which led to reduced fitness in Criollo during the domestication process (Cornejo et al. 2018). The high contribution of Criollo to cultivated cacao helps to explain the low yield on average reported in cultivated cacao globally (Morrissey et al. 2019) and its susceptibility to BPR, WBD and FP diseases (Ploetz 2016). To capture new genetic diversity and to broaden the genetic base for future breeding activities, new plant collections
in northwestern Amazonia should be incorporated into cacao breeding programs (Nieves-Orduña et al. 2023).

In addition, the representation of the first and third clusters in managed cacao likely reflects gene introgression from the Pound Collection. The collection includes accessions MO, IMC, NA, SCA, PA collected in the Peruvian Amazon while searching for trees resistant to WBD (Zhang et al. 2011). The Pound collection has been widely used in cacao breeding for developing disease resistance after the collapse of plantations in Surinam, Trinidad, Ecuador and Brazil due to the introduction of WBD (Zhang et al. 2011). PA series also have been used for developing disease resistance against BPR and FPR (Zhang et al. 2011).

**Genetic resources for cacao breeding**

Guiana (GU) accessions were identified mainly in cluster four in our structure analysis and observed with a very low frequency in managed cacao (Figure 1). The underutilization of Guiana accessions in cultivated cacao was also observed by Cornejo et al. (2018). These findings highlight opportunities to exploit the GU accessions for cacao breeding. Early studies reported cacao trees from French Guiana as novel sources for resistance to BPR and WBD, and high yield (up to 1,426 kg of dry seeds/year/ha) (Lachenaud et al. 2007; Paulin et al. 2008). This was supported by Ofori et al. (2020), who observed that GU accessions can broaden the genetic base of cacao breeding not only for BPR resistance but also yield in Ghana (Ofori et al. 2020). In addition, Guiana accessions in Central America showed moderate resistance to FPR (Lachenaud et al. 2018), an essential agronomic trait for cacao cultivation in Tropical America (Evans et al. 1977; Gutiérrez et al. 2021b; Phillips-Mora et al. 2005, 2013). A detailed agronomic evaluation of multiple Guiana accessions is presented by Lachenaud et al. (2007). The best clones and those to be avoided were identified based on yield, disease resistance (BPR and WBD) and seed quality traits. The preselection of the best Guiana germplasm facilitates
introgression of new and valuable genetic diversity resources into cacao breeding programs (Lachenaud et al. 2007).

**Yield and disease resistance: agronomic traits showing divergence**

Cacao yield traits and disease resistance to WBD showed patterns of divergence between wild and managed cacao accessions (Table 3). These patterns are associated with the history of cacao domestication, cultivation and selection. After initial selection for pulp flavor and seed traits in the sister Curaray population on the Ecuador-Colombia border, human selection led to the domestication of Criollo cacao ~3600 years ago (Clement et al. 2010; Cornejo et al. 2018). In addition, evidence based on ancient cacao DNA supports the consumption of cacao in the Ecuadorian Amazon around 5300 years ago, the DNA analyzed was closer to the Curaray and Purus clusters than to other cacao clusters. (Zarrillo et al. 2018). The genetic cost of cacao domestication led to the accumulation of deleterious mutations, susceptibility to diseases and low yield in Criollo (Cornejo et al. 2018). Criollo with larger seed size, white cotyledons and reduced bitterness was distributed and cultivated outside the Upper Amazon by native Americans in Northern Colombia and Mesoamerica (Cornejo et al. 2018; Motamayor et al. 2002).

Criollo cultivation expanded during colonial times through Tropical America (Bartley 2005), especially in Trinidad, where some blast destroyed the crop in 1727 (Díaz-Valderrama et al. 2020). This collapse in cacao production led to introduction of new plant material from upper Amazonia, which hybridized naturally with the cultivated Criollo and formed a hybrid cultivated cacao known as Trinitario germplasm (Zhang et al. 2011). From these vigorous hybrids, a breeding program was started in 1930 by the Imperial College of Tropical Agriculture of Trinidad focused on yield traits (Toxopeus 1969). The best trees were selected based on the number of seeds per pod and bean weight, which resulted in 100 trees known as the Imperial College Selection (Toxopeus 1969). The mean seed weight was an important trait.
due to the premium price for large seeds in the market (Toxopeus 1969), a trait that still is considered of economic importance in modern cacao breeding programs (Bekele et al. 2022). In addition, after the introduction and impact of WBD in 1932, new wild material with disease-resistance traits was needed in Trinidad. The material searched and collected from the Peruvian Amazon in 1937-1938 includes accessions known as the Pound Collection. This collection created the genetic base for developing breeding resistance against WBD globally (Bartley 2005; Díaz-Valderrama et al. 2020; Evans 2016; Zhang et al. 2011).

In Brazil, the low genetic diversity of cultivated cacao led to the collapse of cacao economies in 1989 due to introduction of WBD in Bahia (Bennett 2003; Evans 2016). In response, breeding programs started in Brazil to broaden the genetic base of cultivated cacao and evaluated germplasm collections for developing resistance against WBD (Bennett 2003).

**SNPs related to flower setting (sexual compatibility)**

Within accessions analyzed here we observed TcSNP1866 with the highest $F_{ST}$ value of 0.260 showing pronounced divergence between wild and managed cacao (Table 3). This SNP was identified in a GWAS study using 295 trees and 5301 SNPs, and incompatibility was measured as frequency of flower retention 15 days after self-pollination (on average 21 flowers were self-pollinated per tree) instead of a yes/no trait (da Silva et al. 2016). Previous studies in cacao highlighted that thousands of individuals are necessary to avoid bias in the estimation of SNP effects associated with small samples sizes (da Silva et al. 2016). In addition, flower dropping is influenced by rainfall, high temperature or insects’ attack which can introduce underestimation in flower retention values (da Silva et al. 2016). Thus, the effect of genotype CC (TcSNP1866) associated with a high percentage (33%) of flower retention may be influenced by small sampling sizes and/or environmentally induced flower dropping (da Silva et al. 2016). From the breeders’ perspective, to select self-compatible trees it is recommended to implement genomic selection which considers thousands of SNPs (da Silva et al. 2016).
Clones common to cacao breeding, including disease resistant and commercial trees are self-incompatible (López et al. 2021; Phillips-Mora et al. 2013). Self-incompatibility in cacao requires plantation designs where cross compatible clones are established to foster the exchange of pollen and field production (López et al. 2021; Phillips-Mora et al. 2013). Since incompatibility is a limiting factor in cacao yield, a common breeding objective is to avoid self-incompatible trees in breeding populations (López et al. 2021).

Although we do not have information on the percentage of flower retention after self-pollinations in the accessions genotyped here that would allow us to validate the phenotypes associated with TcSNP1866, we observed a significantly lower frequency of the genotype CC associated with self-compatibility (da Silva et al. 2016) in managed cacao (Figure 4A), which may reflect the fact that breeding populations and advanced selections include self-incompatible trees. For example, self-compatibility was evaluated in commercial clones such as EET (62, 95, 96, 400), CAUCASIA (37, 39, 43, 47), ICS (1,6,39,60,95), and UF (29, 273, 613, 667, 676 ) (López et al. 2021). These clones (5 to 10 years old) were self-pollinated, and the mean fruit set was 24% across clones, indicating partial self-incompatibility (López et al. 2021). The genotypes (n=18) for these incompatible clones indicated a higher proportion of the homozygous GG (56%), followed by CC (39%) and CG (6%).

As self-compatibility is not absolute in cacao (Lopes et al. 2022), the observation of the genotype CC (TcSNP1866) at higher frequency in wild cacao reflect some degree of self-compatibility in wild populations (Figure 4A). This could be due to geographic origin of wild trees observed isolated along river basins and pollinated by midges with reduced range of movement (Lopes et al. 2022). Accordingly, levels of homozygosity above 70% were observed among wild accessions genotyped with genome-wide SNPs (3K) (Lopes et al. 2022).
SNPs related to yield and pod index

Cacao yield (kg of dry seeds/year/ha) is a polygenic complex trait, at least 40 candidate genes of functional importance encode embryo and seed development, protein synthesis, carbohydrate transport, and lipid biosynthesis and transport (Bekele et al. 2022). In addition, yield components such as number of pods produced per tree, and bean dry weight per pod are influenced by genotypes, environment, agronomic management (e.g., fertilization, shade, irrigation) and diseases (Bartley 2005; Doaré et al. 2020; Fernandes et al. 2020; Phillips-Mora et al. 2013; Solís Bonilla et al. 2022). Disease pressure also influences yield, as high-yielding clones with disease resistance traits exhibited a low diseases incidence (Phillips-Mora et al. 2013). But the same clone can exhibit varying yield potential due to exposition to different pathogen strains (variations in disease pressure) (Jaimes et al. 2011; Jaimes et al. 2019). Disease management in cacao farms also affects the yield potential of clones (Jaimez et al. 2020). For example, removal of FPR-infected pods is a common practice to reduce disease incidence and increase tree productivity (Jaimes et al. 2019; Jaimez et al. 2020).

The SNP Tcm004s00289192 related to increased high yield showed a pronounced divergence between wild and managed cacao (Table 3). Within accessions analyzed here we observed the homozygous genotype GG more frequently in managed cacao (Figure 4A). Among the accessions with the GG genotype are advanced selections such as ICS, UF, TARS and EET clones. The heterozygous genotype AG was observed in high-yielding clones such as TARS, CATIE R6, CCN 51 and VB clones. The allele G (Tcm004s00289192) is associated with a higher yield (Fernandes et al. 2020). Fernandes et al. (2020) identified copy-number variations of SWEET (Sugar Will Eventually be Exported Transporters) genes between the markers Tcm004s00289192 and Tcm004s00615809 on chromosome IV. Likewise, a recent GWAS using a diverse cacao germplasm in Trinidad identified SNPs markers within the genes SWEET17 (chromosome 4) and SWEET2 (chromosome 7) related to cacao yield traits such as...
pod index and seed number (Bekele et al. 2022). Sweet proteins are a family of sugar transporters important for plants biological processes such as growth, development, and response to abiotic and biotic stresses (Singh et al. 2023). SWEET genes in cultivated tree species such as apple (*Malus x domestica*) contribute to fruit sugar accumulation (Zhen et al. 2018), and in *Litchi chinensis* played roles in fruit development, growth and seed development (Xie et al. 2019). SWEET4 also contributed to rice and maize domestication by enhancing seed sizes as it facilitates sugar transport during grain filling (Sosso et al. 2015). Further characterization of cacao SWEET genes in wild and cultivated cacao can help to identify alleles contributing to cacao domestication and validate useful alleles for improving yield traits in cacao.

In cacao breeding pod index is defined as the number of pods (fruits) necessary to produce one kg of dry seeds, a low pod index (14-20) is associated with high yield potential and heavier seeds (Bekele et al. 2020; Fernandes et al. 2020). Early breeding programs in Trinidad (1940s) focused on yield and pod index; selected trees exhibited a low pod index (18) and higher yield (1000kg/ha) (Toxopeus 1969). The pod index also showed a high narrow-sense heritability (0.64) and stability across different sites at two farms in Bahia, Brazil, making the trait a target of selection (DuVal et al. 2017). In addition, a low pod index is also selected in germplasm collections and breeding populations because less healthy fruits are necessary to produce one kg of dry cacao, meaning less costs associated with harvesting and pod breaking (Bekele et al. 2020; Solís Bonilla et al. 2022). The SNP Tcm002s23708704 related to pod index showed divergence between wild and managed cacao (Table 3), and Fernandes et al. (2020) reported the A allele to be associated with high pod index. We observed the genotype AA (SNP Tcm002s23708704) at low frequency in managed cacao compared to wild cacao (Figure 4A). Although we do not have phenotypic information for the accession genotyped here, the low
frequency of the A allele in managed cacao may reflect the continued selection favoring a low pod index in breeding populations and cultivated cacao.

**SNPs related to witches’ broom disease (WBD) resistance**

WBD is a devastating disease for cacao cultivation (Evans 2016). To improve disease resistance, cacao breeders use SCA 6 and SCA 12 clones as primary sources for breeding WBD resistance (Gutiérrez et al. 2016). However, due to the susceptibility of SCA clones under high disease pressure, there was a need to identify new sources of resistance. In this effort, Pereira et al. (2021) identified new clones, such as C SUL-3 and GU-171, resistant to WBD in Bahia, Brazil. In addition, new expeditions to the Peruvian Amazon collected 280 cacao trees to diversify the gene pool and resistance to WBD (Durham 2011). Native to Amazonia but widely distributed in South America, WBD has different pathogen strains (Lisboa et al. 2020; Ploetz 2016). Thus, a major breeding objective is to develop cultivars with broad WBD disease resistance (Meinhardt et al. 2008). WBD resistance is a complex trait involving at least sixteen candidate resistance genes identified using genome-wide association studies (GWAS) (Osorio-Guarín et al. 2020) and quantitative trait locus (QTL) analysis (Chia Wong et al. 2022; Mournet et al. 2020; Royaert et al. 2016).

The SNP Tcm004s00110232 related to WBD resistance showed divergence between wild and managed cacao (Table 3). We inferred the data on reaction to WBD for 92 samples (30%) from the ICGD (Turnbull and Hadley 2023), while 219 (70%) have no data available. The phenotypic data reported are from different WBD studies (Turnbull and Hadley 2023), and due to region specific differences among studies likely responses to different pathogen strains were assessed. The genotypes CT and TT (Tcm004s00110232) are highly represented in resistant (n=58) and tolerant (n=25) plants (Figure 5). The genotype CT observed in the clone TSH 1188 was reported as WBD resistant by Royaert et al. (2016). In addition, the genotype CC (Tcm004s00110232) was observed only in resistant germplasm (Figure 5). These results are in
agreement with Lachenaud et al. (2007) that reported the observed homozygous CC in resistant to WBD Guiana clones (GU 171 /C; GU 219 /F; GU 221 /C; GU 261 /P; GU 277 /G) (Lachenaud et al. 2007; Turnbull and Hadley 2023).

The genotype TT (Tcm004s00110232) was detected in all nine accessions reported as susceptible to WBD (Figure 5) but with more samples for this category with other genotypes could probably be detected. This reaction to WBD is explained by distinct variants of *M. perniciosa* (range of pathogen aggressiveness) and pathogen adaptation to resistance trees such as SCA 6 (Artero et al. 2017; Pereira et al. 2021; Royaert et al. 2016). For example, in Bahia, Brazil, a decrease in resistance of SCA clones to WBD has been reported due to continuous and high disease pressure (Pereira et al. 2021). The fact that the SCA clones have been the main source for breeding WBD resistance motivated the search and evaluation of new sources of disease resistance from different geographical areas (Durham 2011; Pereira et al. 2021). In addition, disease resistance is a polygenic trait with each gene explaining a relatively small portion of the variation in disease resistance.

The majority of wild accessions here genotyped correspond to a subset of samples collected during the 1930-1940s in Peru by Frederick J. Pound (see material and methods), whose main purpose was collecting cacao trees exhibiting WBD disease resistance traits (Bartley 2005; Zhang et al. 2011). This explains why the favorable allele T (Tcm004s00110232) was observed at high frequency in wild cacao (Figure 4B). The high frequency of genotype CT in managed cacao reflect the hybridization of managed cacao with wild cacao accessions. For example, a total of 191 crosses used SCA 6 (low yield but resistant to WBD) as a parent for the incorporation of disease resistance traits in cultivated cacao (Turnbull and Hadley 2023). As a result, commercial hybrids such as ICS, TSH, EET, EQX, and TARS with various levels of WBD resistance were developed using SCA 6 (Turnbull and Hadley 2023).
SNPs related to Ceratocystis wilt (CW) resistance

CW targets the cacao vascular system and causes the death of infected trees (Engelbrecht et al. 2007). The disease is caused by the host-specialized fungus *Ceratocystis cacaofunesta*, which is native to South America (Western Ecuador and Southwest Brazil) (Engelbrecht et al. 2007). The disease is geographically restricted to Tropical America. Still, it threatens the cacao economy because it can be dispersed to important cacao-producing regions such as West Africa and Asia (Engelbrecht et al. 2007). Early reports in the 1950s described CW causing damage to cocoa farms in Colombia, Ecuador, Costa Rica and Trinidad, and in 1997 it was observed in Bahia, Brazil (Cabrera et al. 2016). Breeding for disease resistance is gaining attention as germplasm selected for WBD resistance in Brazil such as “Theobahia” shows susceptibility to CW (Fernandes et al. 2018; Lopes et al. 2011).

The SNP Tcm006s13222057 is associated with resistance to CW (Fernandes et al. 2018). This SNP shows significant differences between wild and cultivated cacao populations (Table 3). To investigate the relationship between SNP genotypes and CW resistance, we searched in the ICGD for the reaction to CW among the accessions genotyped (Turnbull and Hadley 2023). Only 65 accessions (21%) have reported data on the CW reaction (resistant, tolerant or susceptible), while 249 accessions (79%) have no data available. Among the accessions with reported data, we observed that the resistant (n=28) and tolerant (n=16) groups had a high frequency of the G allele (Tcm006s13222057), being the genotypes GT and GG more frequent in the resistant and tolerant group, respectively (Figure 6). While in the susceptible group the genotype TT is prevalent (Figure 6). However, these differences were not significant, likely, due to small sample size. These results are however consistent with the QTL analysis by Fernandes et al. (2018), which identified the G allele as a marker for CW resistance.

We observed the genotype GG at high frequency in wild cacao (Figure 4B). Among these accessions the IMC clones (IMC 11, 31, 47, 60 and 67), PA 121, POUND (12,12A), SCA (6,
12) and U (26, 70) are reported as resistant to CW (Turnbull and Hadley 2023). The wild accession IMC 67 is frequently used in cacao breeding programs for its resistance to CW and vegetative vigor and is widely used as a rootstock (Cabrera et al. 2016; Osorio Montoya et al. 2022). IMC 67 was used in 145 crosses for developing cultivars and breeding populations of the series CEPEC, EET, EQX, TSH, TSH (Turnbull and Hadley 2023). However, IMC 67 was reported to be resistant to the Ecuadorian isolate but susceptible to the Brazilian strain (Cabrera et al. 2016), highlighting the need for new (wild) sources against different CW strains. In addition, we observed the heterozygous genotype GT at high frequency in managed cacao (Figure 4B). Heterozygous clones reported as disease resistant to CW includes EET clones (399, 400), SC 20, ICS clones (6, 40, 95), TSH (1188, 595), UF (29, 650) and VB 650 and VB 681, which is recommended for large scale plating in Bahia, Brazil (Lopes et al. 2011; Turnbull and Hadley 2023).

5 CONCLUSIONS AND RESEARCH QUESTIONS

Accessions structure of wild and managed cacao

Our results confirm a narrow genetic diversity in managed cacao likely reflecting Criollo ancestry of accessions traditionally used to select for chocolate quality. Among the germplasm analyzed, managed cacao (cultivated and breeding populations) showed introgression of wild cacao collected in western Amazonia (Peru, Ecuador), but much less contribution of Guiana clones. New collection trips should be done to broaden the genetic base for cultivated cacao to improve agronomic traits, such as urgently needed FPR disease resistance. Potential areas for cacao germplasm collection are highlighted by Nieves et al. (2023).

New genetic resources for cacao breeding

Guiana accessions represent a genetic resource underutilized in managed cacao. Agronomic evaluations of this germplasm group showed favorable yield traits (e.g., accession GU 285)
and resistance against WBD and BPR (Lachenaud et al. 2007). Cacao breeding programs can obtain Guiana clones free of pathogens through the ICQC and use them in perspective breeding experiments to exploit their potential under a hybridization scheme. Developing of cacao inbred lines can support hybrid breeding but homozygosity and genetic distance between accessions should be considered to maximize heterosis (Akpertey et al. 2022; Lopes et al. 2022).

Validation of SNPs and cacao breeding

Cacao SNPs associated with disease resistance and yield traits should be validated across diverge germplasm added with standard phenotypic information obtained from multi-environment studies and using trees with the same age as well as clonal replicates. This validation facilitates the identification of useful major QTL for cultivar development by different breeding programs. In addition, breeding programs can adopt and optimize genomic selection. Previous studies have demonstrated to improve selection if individual effects of multiple SNP markers are considered in complex cacao traits such as yield and disease resistance (Bekele et al. 2022; McElroy et al. 2018; Romero-Navarro et al. 2017).

Candidate genes of cacao domestication

Disease resistance and yield traits showed divergence between wild and managed cacao, probably reflecting selection during domestication, cultivation and breeding efforts. Further analysis with a diverse natural cacao population, cultivars (e.g., CEPEC’s germplasm of Brazil) and clones developed by chocolate companies for specific farm conditions, using high density SNP genotyping panels (cacao 15K SNP array) will help identify signatures of selection in candidate genes with high phenotypic impact on agronomic traits.
Other traits of economic importance

Plant architecture is a trait of economic importance not widely investigated in breeding programs to improve harvest index (ratio of seeds to total below biomass produced). The breeding challenge is to reduce plant height to facilitate harvesting but without affecting yield. Mustiga et al. (2018) proposed selecting trees with small trunk diameter, lower branch angles, and high yield to increase plant density. In addition, market restrictions in cacao-based products with high cadmium levels have led to evaluating germplasm and identifying valuable genetic sources to introduce low cadmium traits into commercial clones (Lewis et al. 2018).

Supplementary Information The online version contains supplementary material available at

Author contributions HEN-O participated in conceptualization, funding acquisition, investigation, data analysis, figure elaboration, writing original draft, validation, reviewing and editing the manuscript; MM participated in data analysis, validation, reviewing and editing the manuscript; KVK participated in conceptualization, investigation, data analysis, supervision, validation, reviewing and editing the manuscript; OG participated in conceptualization, funding acquisition, investigation, data analysis, project administration, supervision, validation, reviewing and editing the manuscript. All authors commented and edited intermediate versions of the manuscript. All authors reviewed and approved the final manuscript.

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Data availability This research used only published SNP markers, which are summarized in table 1. The genotype data generated within this study is presented in supplementary table S3.

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**Declarations**

**Conflict of interest** The authors declare no competing interests

**Data Archiving Statement** This research used only published SNP markers, which are summarized in table 1. The genotype data generated within this study is presented in supplementary table S3.
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GENERAL DISCUSSION

The results in Chapter 2 demonstrated that cacao has a wide distribution in Amazonia (Chapter 2, Fig. 1). The majority of observation points are concentrated in the northwestern Amazon, the hotspot of cacao genetic diversity (Motamayor et al. 2008). Within this region, an area with low observation of cacao was identified, but expected cacao diversity is high (Zarrillo et al. 2018). This finding suggests that this area holds potential for future cacao collections, as illustrated in Chapter 2, Figure 1. In addition, the potential geographic distribution of cacao in Amazonia suggests that further diversity analysis using new and distinctive wild germplasm will likely reveal new genetic clusters (Cornejo et al. 2018; Zhang et al. 2011). From the observed wide geographic distribution of cacao in Amazonia, only two areas (precipitation ≤ 1000 mm/year) were identified for collecting wild cacao with potential drought tolerance traits (Chapter 2, Fig. 2). These populations should be included in breeding programs in important cacao countries such as Ghana and Ivory Coast where long dry periods are affecting yield (Dzandu et al. 2021; Schroth et al. 2016).

Chapter 2 also shows that fronts of deforestation in Amazonia (Pacheco et al. 2021) are eroding cacao natural habitats, limiting the availability of diverse genetic resources for ex situ conservation efforts (CacaoNet 2012). In addition, cacao expeditions for ex situ conservation focused on searching and collecting trees with disease resistance to WBD (Zhang and Motilal 2016). In this context, Chapter 2 highlights the lack of cacao expeditions targeting wild genotypes exhibiting resistance to the economically important FPR disease (Phillips-Mora et al. 2005).

The review of the genetic basis of agronomic traits such as disease resistance, yield and quality traits (flavor and fat) suggests a polygenic nature (Chapter 2). For example, there are at least 44 genes associated with disease resistance for prospective genetic improvement of cacao (Chapter 2, Table 1). Modern tree breeding programs, instead of relying on major QTL, have
adopted genomic selection, that can predict complex traits using thousands of genome-wide SNP markers, accelerating breeding cycles and increasing selection intensity (Grattapaglia et al. 2018; Grattapaglia 2022). In cacao, genomic selection has demonstrated to improve selection since individual effects of multiple SNP markers are considered in complex cacao traits such as yield and disease resistance (Bekele et al. 2022; McElroy et al. 2018; Romero-Navarro et al. 2017).

The high chloroplast diversity observed in northwestern Amazonia highlights this area as a center of cacao genetic diversity (Chapter 3). The observation of 23 chloroplast haplotypes including seven geographically restricted haplotypes suggests that in situ conservation areas can be proposed in northwestern Amazon (Chapter 3, Fig 2-3). Previous studies of tropical tree species such as Cedar (Cedrella odorata) used the geographic distribution of chloroplast haplotypes to propose areas for in situ conservation in Central America (Cavers et al. 2003). The pattern of dispersal of chloroplast haplotypes observed from the west to east Amazon suggests a decline in cacao genetic diversity (Chapter 3, Fig 2-3). This geographic pattern of decline in cacao genetic diversity was also observed by Cornejo et al. (2018). This could be explained by past human influences, as cacao was spread in Amazonia by humans, as it was the case with other tropical tree species with crop potential (Clement 1999; Levis et al. 2017).

The structure analysis of cacao germplasm using SNPs associated with agronomic traits identified four genetic clusters reflecting the geography origin of samples (Chapter 4, Figure 1). Managed cacao (breeding populations and cultivated varieties) was represented mainly by two genetic clusters, suggesting a low population structure. This germplasm structure supports the findings of Cornejo et al. (2018). According to their research, modern man-made cacao hybrids are largely represented by two out of ten identified cacao genetic clusters (Cornejo et al. 2018; Motamayor et al. 2008). These results suggest the opportunity to broaden the genetic base of cultivated cacao which is hampered by diseases and low yield (Bekele and Phillips-
Mora 2019; Gutiérrez et al. 2016). For example, Guiana germplasm was observed in low proportion in managed cacao but previous agronomic evaluations suggest that disease resistance and yield traits can be exploited by cacao breeding programs (Lachenaud et al. 2007). Divergence of wild and managed cacao ($F_{ST} > 0.05$) suggest selection processes in cacao germplasm in agronomic traits such disease resistance (WBD and CW) and yield (pod index and high yield) (Chapter 4, Table 3). These patterns are associated with historical process in cacao selection and cultivation (Toxopeus 1969; Zhang et al. 2011)

The heterozygous genotypes observed in cacao accessions with disease resistance traits suggest QTL that potentially contribute to cacao heterosis. For example, the genotypes CT (SNP Tcm004s00110232) related to witches’ broom disease (Royaert et al. 2016) and the genotype GT (SNP Tcm006s13222057) related to Ceratosystis disease (Fernandes et al. 2018) were observed in accessions reported as disease resistant-tolerant (Chapter 4, Fig 5-6). Validation of SNP markers associated with agronomic traits should be validated for different cacao breeding programs to facilitate the identification QTL for cultivar development.
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CONCLUSIONS

The conservation and use of cacao genetic diversity in breeding programs are vital for sustainable cacao cultivation and the global chocolate industry. The geographic distribution of cacao pinpoints to new areas for germplasm collection and supports northwestern Amazon as the hotspot of cacao genetic diversity. This was also validated by the observation of areas with high haplotype diversity in the Peruvian Amazon. In addition, observations of wild cacao in Amazonia indicate the existence of natural populations that have not yet been subjected to population genomic analysis, and have not been included in breeding efforts. New genetic clusters with allelic variants that influence agronomic traits can be identified. However, trends in Amazon deforestation limit the conservation and use of cacao primary gene pool.

Cacao germplasm (cultivated and breeding populations) shows a narrow genetic diversity reflecting historical processes of selection. Agronomic traits that distinguish wild and managed cacao include disease resistance to WBD and yield traits. Germplasm resources observed with low contribution to managed cacao but with agronomic potential include the Guiana germplasm. Cacao SNPs linked to agronomic traits need to be validated under different environments to facilitates the identification of major QTL for cultivar development.

Overall, cultivated cacao can benefit from novel genetic resources available in wild germplasm or obtained from new collections in northwestern Amazonia. Standardized agronomic and genotypic evaluation add value to cacao genetic resources held in ex situ collections and improve its utilization in breeding programs. Challenges remain for the pyramiding of agronomic traits in cacao, such as developing inbred lines to exploit heterosis and genomic selection in important agronomic traits such as disease resistance to BP, WBD and FPR, yield and seed quality traits.
OUTLOOK

Based on the results presented in chapters 2, 3, and 4, future research areas within the frame of cacao genetic diversity and agronomic traits that will contribute to building breeding populations and ultimately developing cultivars are presented.

Population differentiation studies using neutral molecular markers are likely to reveal new cacao clusters, and forest areas in northwestern Amazonia with potential for collecting diverse germplasm are presented in Fig. 1, Chapter 2. In addition, collecting and conserving diverse cacao germplasm from wild habitats will also increase the availability of genetic resources for breeding and help to understand how cacao genetic diversity varies geographically and is distributed across Amazonia. In this context, further analysis with more diverse samples and informative cacao chloroplast SSR markers (Chapter 3) will help to better understand the phylogeographic structure of cacao in Amazonia. Unfortunately, the natural habitat of cacao is disappearing due to deforestation (Fig. 1, Chapter 2), hindering future research progress in population genetic analysis and cacao germplasm conservation.

Cacao germplasm conserved in *ex situ* collections should be analyzed with SNP markers to identify genetic gaps, mislabeling and redundancy, followed by standard agronomic evaluations to establish an efficient conservation and increased use of cacao genetic diversity in cultivar development (Chapter 2). These analyses are critical to increase the genetic diversity of cultivated cacao, as we observed two overrepresented genetic clusters among cultivars and breeding populations (Chapter 4). Novel genetic resources are needed to broaden the genetic base of breeding populations in support of agronomic traits. For example, the agronomic potential of Guiana germplasm observed in less proportion in breeding and cultivated cacao should be evaluated under different environments (Chapter 4).

Research on traits of economic importance includes identification of novel alleles associated with resistance to different pathogen strains of WBD and FPR (Chapter 2). These
studies will benefit by using diverse cacao germplasm (Chapter 4) and standard phenotyping allowing comparability of genotype performance between breeding programs.

Introgressions of quality traits in cacao such as low seed Cd accumulation are needed in cacao cultivars (Chapter 2). Identification of genotypes with low seed Cd accumulation are needed to select potential rootstocks. The breeding challenge is to combine low Cd with disease resistance traits in cacao germplasm used as rootstock (e.g., IMC 67). Cacao candidate genes associated with Cd transportation for perspective genetic improvement of cacao include \textit{TcNRAMP5}, \textit{TcHMA3}, and \textit{TcHMA2}. Thanks to advances in cacao gene editing, these genes can be targeted to reduce seed Cd accumulation. Still, it requires high-quality reference genomes and protocols for somatic embryogenesis in cacao cultivars (Chapter 2).

Drought tolerance is a trait not well studied in cacao breeding programs. The identified natural populations growing in areas with precipitations \(\leq 1000\) mm/year (Fig. 2, Chapter 2) can help to build breeding populations to increase drought tolerance by targeting root traits such as angle and root biomass that improve water use efficiency.

Although there are advances in developing major QTL for cacao breeding (Chapter 4), future research areas in cacao breeding should implement genomic selection. For example, development of cacao inbred lines to exploit heterosis followed by genomic selection of untested clones under different environments should be performed. Genomic resources such as the cacao Infinium array (15K SNPs) should be used by breeding programs in analytical and experimental studies such as population differentiation, association studies and genomic selection.
SUMMARY

The tropical cacao tree, *Theobroma cacao* L., is cultivated to produce seeds, the unique raw material for the chocolate industry. Thus, conservation and use of cacao genetic resources in breeding programs are vital for the cacao-chocolate global economy. Wild cacao populations have a wide geographic distribution in their native range along the Amazon basin, but northwestern Amazonia is the hotspot of cacao genetic diversity. From the rich diversity of cacao (at least twelve genetic clusters are identified) the domestication of cacao in South America led to the development of Criollo, whose seeds are characterized by reduced bitterness, a trait appreciated for the international cacao market. Criollo ancestry still is a primary source for the development of cultivated cacao varieties. A narrow genetic base in cultivated cacao and disease vulnerability led to collecting disease resistant genotypes in Amazonia since the 1930s. This was followed by conservation via *ex situ* collections, agronomic evaluation of accessions to improve and broaden the genetic base of cultivated cacao. Disease resistance and yield are the most important economic traits in cocoa breeding, as diseases can destroy up to 40% of cocoa production. The conservation and utilization of the rich cacao diversity are the primary source of novel alleles necessary to address current and future cacao breeding objectives.

The first paper reviews key topics in cacao genetics essential for crop improvement. By analyzing the geographic distribution of wild cacao in Amazonia, which consisted of 4413 observation points compiled from the years 1938 to 2022, and mapping the patterns of genetic variation, new areas for germplasm collection were identified in the border region between Ecuador and Peru in northwestern Amazonia. Despite the deforestation in Amazonia, this analysis suggests that it is still possible to collect wild material in the identified areas. In terms of conservation, germplasm collections in cacao producing countries can benefit from available
SNP markers to improve genotype identity and characterization of agronomic traits at early stages.

The studies addressing the genetic basis of disease resistance, seed quality traits, and yield suggest a polygenic architecture. For example, at least 44 genes reported and distributed over the ten cacao chromosomes are associated with resistance to three different diseases (Black Pod-BP, Witches’ Broom Disease WBD and Frosty Pod Rot-FPR). Further evaluations with more diverse germplasm and different fungal strains likely will reveal more disease resistance genes and genotypes. Recently, the genetic basis of seed quality traits such as reduced cadmium and flavor has been addressed, but informative molecular markers need to be developed to accelerate breeding. Other traits of particular importance to West Africa, the largest producer of cacao, include heat and drought tolerance. Cacao breeding efforts have not fully exploited heterosis but can profit from modern genome-wide SNP data to develop inbred lines and design crosses, followed by genomic selection of untested hybrids. Overall, cacao breeding requires data generated by standard phenotyping (of different cacao populations) integrated with analytical methods such as genome-wide association studies (GWAS), quantitative trait locus (QTL) mapping, genomic selection, and field trials to identify relevant genotypes for breeding and cultivation.

The second paper investigates the phylogeographic structure of cacao using nine informative chloroplast DNA (cpDNA) microsatellite (simple sequence repeat - SSR) markers to genotype 233 cacao samples. Nineteen out of the twenty-three chloroplast haplotypes identified were observed in western Amazonia, including seven geographically restricted haplotypes, confirming northwestern Amazonia as a hotspot for cacao genetic diversity and center of crop domestication. Two observed haplotypes were widespread from west to east along the Amazon basin, suggesting seed dispersal by humans. When comparing the observed haplotypes with previously identified genetic clusters using nuclear SSRs, specific chloroplast
haplotypes were observed within eight genetic clusters. Evidence of selection of plant material was provided by the observation of a single haplotype common in cacao farms of west Ecuador and reference Trinitario accessions, the most important breeding population. The identified chloroplast haplotypes can be used to determine the geographic origin of planting material at a finer geographic scale. In addition, these chloroplast haplotypes, in combination with phenotypic assessments, can help to select geographically distinctive varieties for cacao breeding programs.

The third paper analyses the population structure of wild and managed cacao (cultivated and breeding populations), identifies agronomic traits under selection, and uncovers valuable cacao genotypes not observed in cultivated and breeding populations. Forty published SNPs associated with disease resistance and 11 SNPs with yield traits were investigated in 346 accessions using the MassARRAY® system. The structure analysis revealed four genetic clusters. Two clusters were overrepresented in managed cacao, which is characterized by admixed individuals, representing artificial hybrids. A cluster overrepresented in managed cacao is linked to the preference for maintaining the superior flavor of cultivated cacao from the Criollo ancestry. Under-utilization of genetic resources such as Guiana cacao was detected in managed cacao, and published agronomic evaluation of this germplasm group showed yield and disease resistance traits. Yield and disease resistance traits (mainly resistance to WBD) show divergence between wild and cultivated cacao ($F_{ST} > 0.05$). This divergence likely reflects selection during domestication, cultivation, and breeding. The identified SNPs showing divergence between wild and cultivated cocoa can be used to improve selection in breeding populations. Agronomic evaluation of wild populations can expand the genetic base for breeding cacao, especially for traits of economic importance such as low seed Cadmium and disease resistance against FPR.
ZUSAMMENFASSUNG


Der erste Artikel befasst sich mit Schlüsselthemen der Kakaogenetik, die für die Verbesserung von Nutzpflanzen unerlässlich sind. Durch die Analyse der geografischen Verbreitung von Wildkakao in Amazonien, die aus Beobachtungspunkten aus den Jahren 1938 bis 2022 bestand, und die Kartierung der Muster der genetischen Variation wurden neue
genomischer Selektion und Feldversuchen zur Identifizierung relevanter Genotypen für Züchtung und Anbau kombiniert.
