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Traditional agroecosystems as conservatories and incubators of cultivated plant varietal diversity: the case of fig (Ficus carica L.) in Morocco

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Abstract

Background: Traditional agroecosystems are known to host both large crop species diversity and high within crop genetic diversity. In a context of global change, this diversity may be needed to feed the world. Are these agroecosystems museums (i.e. large core collections) or cradles of diversity? We investigated this question for a clonally propagated plant, fig (Ficus carica), within its native range, in Morocco, but as far away as possible from supposed centers of domestication.

Results: Fig varieties were locally numerous. They were found to be mainly highly local and corresponded to clones propagated vegetatively. Nevertheless these clones were often sufficiently old to have accumulated somatic mutations for selected traits (fig skin color) and at neutral loci (microsatellite markers). Further the pattern of spatial genetic structure was similar to the pattern expected in natural population for a mutation/drift/migration model at equilibrium, with homogeneous levels of local genetic diversity throughout Moroccan traditional agroecosystems.

Conclusions: We conclude that traditional agroecosystems constitute active incubators of varietal diversity even for clonally propagated crop species, and even when varieties correspond to clones that are often old. As only female fig is cultivated, wild fig and cultivated fig probably constitute a single evolutionary unit within these traditional agroecosystems. Core collections, however useful, are museums and hence cannot serve the same functions as traditional agroecosystems.

Background

High yield agriculture based on elite crop varieties and high inputs results in loss of both numbers of crop plants and genetic resources within crops, thus threatening crop biodiversity and the nutritional safety of humanity [1]. To preserve crop diversity, traditional landscapes may have to be preserved [2]. In analogy with the concept of “biodiversity hotspot” used to identify priority areas for the conservation of wild species [3], traditional agroecosystems could be considered as main conservatories of crop biodiversity [4]. Indeed in 2002 the FAO started an initiative for the conservation and adaptive management of Globally Important Agricultural Heritage Systems http://www.fao.org/nt/giahs/en/. Although they are quite diverse, these agroecosystems exhibit common features such as 1) a high diversity of crop species, 2) the use of diversified traditional varieties, 3) sustainable agriculture, 4) low inputs associated with traditional farming practices and 5) the farmers obtaining a sizable proportion of their seeds (or vegetative equivalents) from their own harvest [5]. For instance, a survey in continental oases in northern Oman recorded 107 different crop species belonging to 39 families, including 33 fruit species [6]. This large biodiversity was successfully achieved despite the constraints of a small scale cropping system under arid and semi-arid conditions. Similarly, a study of 27 crop species in traditional agroecosystems distributed in eight countries over the five continents [7] demonstrated that such agroecosystems maintain considerable within crop genetic diversity. Traditional agroecosystems are either the repositories of crop diversity, or the place where extant crop diversity was fostered. Hence investigating within crop species genetic diversity and its geographic variation would help understanding genetic resources.
and dynamic processes of past and present domestica-
tion and subsequent diversification.

The biodiversity hotspot concept is associated with a
major pattern of biodiversity: it increases close to the
equator, and decreases towards the poles [8]. Two main
ideas have been suggested to explain this global biodi-
versity pattern. Equatorial regions are a museum of bio-
diversity preserving ancient biodiversity, and/or they are
a cradle generating new biodiversity [9]. If agroecosys-
tems are hosting huge crop biodiversity, should we con-
sider them as museums or as incubators of crop
biodiversity, or as both? For long term crop manage-
ment policies and hence to feed the world, the answers
to this question is of a great importance.

The Mediterranean basin is one of 25 hotspots of bio-
diversity in the world. It hosts 25,000 species, of which
13,000 are endemic, this later group representing 4.3%
of the worldwide flora [3]. It is the largest biodiversity
hotspot on earth (over 2,000,000 km²) and it includes
several separate refuge areas [10]. Traditional agroeco-
systems are still found all over the Mediterranean region
in mountains and oases. However several of these tradi-
tional agroecosystems may be of particular importance
for preserving crop biodiversity. Indeed, many plant spe-
cies were originally domesticated close to the Eastern
shores of the Mediterranean. Hence, we might encour-
gage contrasted patterns of genetic diversity within crops
together the Mediterranean region, with more crop
diversity available in the Eastern Mediterranean.

The process of domestication seems to have been dif-
suse, with prolonged cultivation of undomesticated forms,
and prolonged genetic exchanges of domesticated forms
with local wild relatives, at least for crops propagated by
seeds [11,12]. With a such domestication process, tradi-
tional agroecosystems located in the East Mediterranean
may be most important for preservation of crop genetic
resources. In addition, the domestication process of clon-
ally propagated crops, particularly fruit trees, is often
thought to have been an instant or almost instant process
[13,14] building on the idea that genotypes presenting the
whole suite of agronomic traits of interest may have arisen
by chance within totally natural populations [15]. This
may qualify as a silver bullet hypothesis. If we follow this
hypothesis, domestication was instantaneous, and followed
by subsequent clonal propagation. Hence we would expect
that extant varieties are old, probably limited in number,
and that they represent the gene pool of the original
region of domestication. The wild progenitors of some of
these clonal crops still grow all around the Mediterranean
region. This is true for three most symbolic crops in these
regions such as olive, grape wine and fig. Therefore, we
may ask, within such species, whether extant varietal
genetic diversity in traditional agroecosystems reflects the
propagation of old widespread clones, or old local clones,
or recent local clones. We may even ask whether varieties
could be fuzzy aggregations of genotypes (landraces) [16].

We chose to address this question in fig which pre-
sents us with a particularly fascinating situation as it is
extremely easy propagated via cuttings, and was domes-
ticated extremely early in the Near East, contemporarily
with cereal crops, 9-12,000 BP [17]. Fig, *Ficus carica* L.,
is dioecious. Female trees produce the edible crop. Male
trees produce pollen and their figs host the pollinator,
*Blastophaga psenes* [18]. Each fig variety is a clone of
female tree that are propagated through cuttings. Some
fig varieties may produce seedless fig fruits without pol-
lination while other varieties require pollination for suc-
cessful fruit set [19]. Female figs produce seeds if
pollinated. Male figs are often collected far from zones
of fig cultivation and suspended in cultivated female
trees to ensure pollination [20].

Phylogeographic studies based on cytoplasmic genes
showed that wild fig was present all over the Medi terr-
anean basin before domestication [21]. We investigated
the genetic diversity of fig varieties in Moroccan tradi-
tional agroecosystems. Morocco is at the Western limit
of the natural range of fig, as far away as possible (over
3500 km) from postulated places of domestication.
Hence, if domestication begun and ended in the Eastern
Mediterranean, then we expect to observe limited diver-
sity so far away from the original zone of domestication.
We also expect to observe lack of spatial genetic struc-
ture within Morocco, or simply a decrease of diversity
when further away from the shores of the Mediterranean.

We made extensive collections of fig cultivars *in situ*,
in order to 1) test whether cultivars are effectively highly
local, 2) detect whether some of these cultivars are old,
and 3) establish what insights into the history of fig cul-
tivation could be drawn from extant genetic diversity
and its spatial structuring.

We show here that in traditional agroecosystems, fig
varieties are true clones, highly diversified, often highly
local. Nevertheless they are often sufficiently old to have
accumulated somatic mutations. Spatial genetic struc-
ture resembles what would be expected for a wild plant
at mutation/drift/migration equilibrium. We conclude
that the Moroccan traditional agroecosystems are at the
same time museums and incubators of fig variety diver-
sity, in a dynamic system preserving old, local varieties
and generating new ones locally.

**Results and Discussion**

277 cultivated trees were sampled throughout traditional
Moroccan agroecosystems distributed over 40 sites that
we grouped into 6 geographical zones (Figure 1). During
field collection, we noted that, within each site, trees
designated by the same name (local variety) shared
highly similar morphological traits. To maximize genetic
diversity of our sampling we generally collected a single individual per variety per site. Nevertheless, in a number of cases we sampled twice the same local variety within a site or within adjacent sites. Such samples systematically shared a same genotype. Hence genetic evidence confirms the obvious conclusion from phenotypic observation that local varieties are generally clones.

SSR polymorphism and its discrimination power
The 277 individuals genotyped were separated into 194 distinct molecular profiles using 17 SSR loci (see Additional File 1). Genetic parameters for each locus are given in Table 1[22-25]. Overall, observed heterozygosity was higher than expected heterozygosity. The discriminating power per locus, $D_i$ (probability of distinguishing two randomly chosen clones), ranged from 0.495 (LMFC26) to 0.979 (LMFC30) with a mean of 0.70 (Table 1). Hence the probability of confusing a randomly chosen clone with another one (under the hypothesis of statistical independence of the loci) was $I(1-D_i) = 5 \times 10^{-11}$. With only 38,226 pairwise comparisons (including identical genotypes) in our data set, all cases of genotype identity should correspond to clones.

Table 1 Genetic parameters of the 17 SSR loci used in this study

| Locus | $A$ | Size range (in bp) | $H_O$ | $H_E$ | $F_{IS}$ | $D$  |
|-------|-----|--------------------|------|-------|---------|------|
| MFC1a | 5   | 161-195            | 0.620| 0.629 | 0.016   | 0.841|
| MFC2a | 7   | 156-190            | 0.599| 0.602 | 0.008   | 0.880|
| MFC3a | 9   | 96-136             | 0.818| 0.760 | -0.074  | 0.851|
| MFC4a | 5   | 216-226            | 0.524| 0.493 | -0.060  | 0.652|
| MFC8b | 2   | 173-177            | 0.508| 0.490 | -0.033  | 0.619|
| MFC9b | 7   | 188-211            | 0.636| 0.582 | -0.090  | 0.786|
| MFC11b| 7   | 181-203            | 0.604| 0.569 | -0.059  | 0.585|
| MFC12b| 4   | 152-167            | 0.578| 0.552 | -0.045  | 0.743|
| FSYC01c| 5   | 117-160            | 0.455| 0.451 | -0.005  | 0.842|
| FSYC04c| 2   | 181-183            | 0.529| 0.502 | -0.053  | 0.595|
| LMFC19d| 8   | 296-312            | 0.433| 0.398 | -0.086  | 0.573|
| LMFC24d| 4   | 272-278            | 0.460| 0.456 | -0.006  | 0.646|
| LMFC26d| 3   | 224-236            | 0.235| 0.223 | -0.051  | 0.495|
| LMFC28d| 5   | 192-203            | 0.562| 0.558 | -0.004  | 0.733|
| LMFC30d| 11  | 231-261            | 0.904| 0.820 | -0.100  | 0.979|
| LMFC32d| 9   | 197-225            | 0.433| 0.415 | -0.039  | 0.628|
| LMFC33d| 2   | 245-247            | 0.492| 0.486 | -0.009  | 0.519|

A: number of alleles, $H_O$: observed heterozygosity, $H_E$: expected heterozygosity, $F_{IS}$: within population fixation index, $D$: discriminating power. Primers developed by: a Khadari et al [22], b Achtak et al [23], c Ahmed et al [24] and d Giraldo et al [25].
We plotted the distribution of number of allelic differences between the 194 different genotypes in order to visualize the distribution of genetic differences between genotypes, (Figure 2A; 18,721 pairwise comparisons, excluding identical genotypes). The distribution ranged from 1 to 34 differences, presented a major peak at 19-20 differences and a very distinct, but very small, peak at 1-3 differences. The probability to observe by chance two or more genotypes that were distinguished by 3 alleles was $2.6 \times 10^{-6}$. Further, individuals whose genotypes were identical or differed by only 1-3 alleles were morphologically highly similar (see Additional File 2). The systematic association of genetic similitude for neutral markers with morphological similarity allows to conclude that all these trees belonged to a single original clone and that some had accumulated somatic mutations. Further, the shape of the pairwise genetic difference curve suggests that, beyond the case of the

Figure 2 Frequency distribution of genetic dissimilarity for all pairwise comparisons between cultivated fig genotypes. (A) complete data set; (B) in mountain agroecosystem; (C) in oasis agroecosystem. Genetic differences among genotypes are retained in the oasis agroecosystem, despite the low number of genotypes cultivated (21). Note on the three graphs the bimodal shape of the curve with a very small peak for differences of 1-3 alleles.
few genotypes deriving from each other by somatic
mutations, all other genotypes are the product of sexual
reproduction. We chose to be highly conservative in our
estimate of which genotypes represented somatic muta-
tions. Indeed the curve suggests that the limit may be
better placed above 6 differences and indeed the prob-
ability of observing by chance two genotypes differing
only by 6 alleles was still low, at 0.0017.

Hence, we classified the 194 genotypes into 152 gen-
type groups (clones) separated by at most 3 alleles,
which were distinguished from all other genotypes by
4 to 34 alleles. Out of these groups of genotypes, 128
contained a single individual while 24 groups con-
tained more than one individual and represented collect-
ively 66 genotypes. Often a variety name was found
to be associated with the same clone (identical or
almost identical genotype) in different sites, conforting
our conclusions. Numbers of trees sharing the same
genotypes are given in Additional File 3, while Addi-
tional File 4 and Figure 3 provide a series of cases of
mutations in somatic lines, the presence of such muta-
tions within clonal lineages suggests that these
varieties are old.

Variety names and characterization
Out of 277 sampled fig trees, 246 were named by the
local farmers while for 31 fig trees, the interviewed
farmers did not provide any name (see Additional File
1). These 31 unnamed trees corresponded to 30 geno-
types out of which four corresponded genetically and
morphologically to known varieties ('Ikoran Imelalen',
'El Messari', 'El kehla' and 'Beyota') and three were very
similar genetically and morphologically to the 'Saaidi'
and 'Rhoudane' varieties. The remaining 23 genotypes
were distinct from previously defined varieties.

Synonymy was observed for 23 genotypes, with 2 to 7
denominations per genotype (Figure 3, Additional File
3). Two situations were observed. True synonymy was
observed when the different fig trees presented identical
pomological traits such as the varieties 'Johri' and 'El
Messari' (green fig skin color, flattened pyriform fruit
shape and red internal color). This situation was
encountered for 20 genotypes. False synonymy was
observed for fig trees known under the same generic
denomination to which a descriptor of fig skin color
was added. In these cases the leaves and the figs pre-
sented similar morphologies but fruit color was differ-
ent. Six instances of the latter situation were
encountered (Figure 3, Additional File 3). They included
for instance 'Saaidi Lbyed-IB5-T4-P014' (white skin
color) and 'Saaidi Lkhel-IB5-T3-P014' (black) in the
North west zone, 'Ikoran Ihebchan-IVA2-T2-P002'
(black) and 'Ikoran Imelalen-IVA1-T1-P001' (white) in
the Center zone (Figures 1 and 3).

We suggest that the second type of synonymy corre-
sponds to cases of somatic mutations. A similar situa-
tion has previously been reported in Cataluña for 'Col
de Dame blanche', 'Col de Dame grise' and 'Col de
Dame noire' which are genetically and morphologically
identical and only differ by skin color [26] and in Slove-
nia for 'Green Matalon' and 'Black Matalon' [27]. Such
mutations have been reported in Vitis vinifera [28], and
indeed, in Brazil, a single wine producer successfully
selected 2 clonal color variants [29]. In our study, each
time we encountered several color forms within a vari-
ety, they occurred within the same zone, but not neces-
sarily within the same site (see Figure 1 and Additional
File 3). This suggests that varieties have a prolonged
local history.

We grouped several variety names as highly similar
because they had the same meaning albeit in different
languages or dialects (see Additional File 4). For
instance, the names 'Ikoran Ihebchan -IVA1-T3P041',
'Kahla-VIA1-T4P177', 'Kohli-IA2-T8P018', and
'Taberchante-VA1-T1-P077' sampled in the central
region, in the oases, in the North west and in the Mou-
louya valley, respectively, all corresponded to black figs
presenting turbinate fruit shape, but their genotypes
were distinctive. Thus cases of homonymy involved 31
distinct denominations corresponding to 181 fig trees
and 147 genotypes (see Additional File 4). In a number
of cases such homonymy corresponded to highly similar
genotypes. Nevertheless, the denominations representing
most cases of homonymy were referring to fruit color.
Denominations referring to White, Black and Green
color represent a total of 55 genotypes, i.e. 1/3 of the
164 genotypes sampled with variety denomination.

Depending on the genetic relationships between geno-
types, three types of homonymy were distinguished (see
Additional File 4). First we observed homonymy
between highly similar genotypes (= within a clone)
such as within the varieties 'Rhoudane', 'Zerki' and 'Byed',
which included respectively three, four, and four
very closely related genotypes. As stated above these
correspond most probably to cases of somatic mutation
within clone, and do not really constitute cases of
homonymy. Second we observed cases of homonymy
grouping varieties presenting similar pomological traits
but clearly distinct genotypes, such as the cultivars 'Aïn
Hajla', 'Rhoudane', 'Kehla' and 'Biyadi', representing
respectively two, six, eight, and nine distinct geno-
types. Finally we observed cases of homonymy grouping
varieties presenting different pomological traits and dif-
ferent genotypes (six cases; Additional File 4).

Only eight clones were present in several geographic
zones. This was the case for instance for 'Assel-IA1-
Figure 3 Genetic similitude among fig varieties. Samples grouped within a box correspond to highly similar genotypes that most probably derive from each other by somatic mutation. These similar genotypes often bear similar variety names. After the variety name, the roman number indicates zone of sampling, the letter the subzone, followed by a number giving the precise site of sampling, Tx indicates the tree number x within site and Pxxx indicates genotype number xxx (see Additional File 1).
T4-P010’ (North west zone), ‘Assal-IID1-T6-P010’ (Rif zone), ‘Zerka-VA2-T1-P010’ (Moulayouya valley). These eight non local clones corresponded to widely known varieties, such as ‘Assal-IID1-T6-P010’ = ‘Sebtawi-IA1-T1-P010’; ‘Rhoudane-IIF1-T12=P006’ = ‘Rhoudani-IIIA1-T4-P006’ = ‘El Kehla (Rhoudani)-VC1-T1-P006’ and ‘Bacora-IA3-T5-P019’ = ‘Lemdar-IIIC1-T5-P019’ (see Additional File 3). Hence, in Morocco, most fig varieties are cultivated over a limited spatial. Concurrently, within a geographical zone, varieties often correspond to a single specific clone. For instance, in the Rif, the 81 trees analyzed were assigned to 43 named varieties (and 7 unnamed) and corresponded to 64 genotypes (grouped into 35 clones when including within a clones all genotypes that differed by at most three alleles).

Hence in traditional Moroccan agroecosystems fig local varieties are clones and they are generally highly local and diversified (on average 8 local varieties were collected per site in the Rif region). At least some of these local varieties were sufficiently old to have accumulated somatic mutations on neutral genetic markers and on selected traits.

**Genetic diversity within and among geographical groups**

Similar numbers of alleles were observed within each geographic zone, except the North center zone which presented fewer varieties, few local genotypes and as a consequence fewer alleles (Table 2). Surprisingly in the South zone, all genotypes were local and allele diversity was similar to that observed in other zones. Among the 95 observed alleles, three were exclusively detected in the center zone (MFC3-133, LMFC30-259, LMFC28-192), two in the Moulayouya valley (MFC9-188, LMFC24-278), two in the North west zone (LMFC19-306, LMFC32-225) and four in the South zone (MFC3-96, MFC2-190, MFC9-211, LMFC30-243). Expected heterozygosity was highest in the South zone (0.558) and lowest in the Rif zone (0.495).

There is no published data available on fig genetic diversity in traditional agroecosystems based on a sufficient number of genetic markers to discriminate clones. However, ongoing work in Lebanon and in the Tizi Ouzou area (Algeria) using the same markers (Chalak, *pers. comm.;* Daoudi, *pers. comm.*) suggest the presence of similar level of diversity as in Northern Morocco. These areas correspond to traditional agroecosystems mainly based on subsistence agriculture, with orchards presenting several fruit species grown together and several varieties per species [30,31]. Hence, the pattern observed for fig variety diversity in Morocco can probably be transcribed to most traditional agroecosystems around the Mediterranean. How the pattern may shift outside the range of wild *Ficus carica* remains an open question.

Genetic differentiation among the six geographic zones was about 4% ($F_{ST} = 0.038$). Pairwise comparisons showed contrasted $F_{ST}$ values ranging from 0.017 to 0.068 (Table 3). The highest differentiation ($F_{ST} = 0.07$) was noted between the Southern zone and the Rif zone. These two zones were also the sole zones clearly separated on the two first coordinate axes of the Factorial Correspondence Analysis (Figure 4). A significant spatial genetic structure was observed ($p < 10^{-6}$). Pairwise Loiselle kinship coefficients decreased significantly with distance (Figure 5), and were more strongly correlated with log than with linear distance, whatever the range of distances incorporated in the calculus. Such a pattern would be interpreted in natural populations as isolation by distance with no rupture in gene flow [32].

We may reconcile the three sets of analyses (FCA, $F_{ST}$ and pairwise Loiselle kinship coefficients) by suggesting that we have here the image of spatial genetic structure as could be expected in natural populations for a situation of mutation/migration/ drift processes at equilibrium resulting in some geographic variation in genetic background without geographic variation in genetic diversity. Within this global pattern, the North west

| Geographic zone | trees analyzed | named varieties | unnamed varieties | genotypes | local genotypes | $N$ | $N_a$ | $H_e$ | $H_o$ | $F_{st}$ | p-value |
|-----------------|----------------|-----------------|------------------|-----------|----------------|-----|--------|------|------|--------|--------|
| North west (I)  | 60             | 43              | 4                | 36        | 31             | 66  | 3.88   | 0.523| 0.559| -0.0699| 0.0187 |
| Rif (II)        | 81             | 43              | 7                | 64        | 58             | 70  | 4.12   | 0.495| 0.540| -0.0968| 0.0003 |
| Mountain agroecosystems | 141 | 76              | 11               | 96        | 89             | 77  | 4.53   | 0.510| 0.548| -0.0724| 0.0005 |
| North center (III) | 20            | 12              | 4                | 15        | 12             | 54  | 3.18   | 0.533| 0.558| -0.0502| 0.1477 |
| Center (IV)     | 61             | 23              | 10               | 45        | 40             | 70  | 4.12   | 0.511| 0.557| -0.0904| 0.0156 |
| Moulouya valley (V) | 34             | 19              | 2                | 27        | 24             | 70  | 4.12   | 0.518| 0.507| 0.0219| 0.4292 |
| South (VI)      | 21             | 15              | 3                | 21        | 21             | 72  | 4.24   | 0.558| 0.571| -0.0242| 0.2898 |
| Oasis agroecosystems | 21            | 14              | 3                | 21        | 21             | 72  | 4.24   | 0.558| 0.571| -0.0242| 0.2898 |

$N$: total number of alleles observed within each zone; $N_a$: mean number of alleles per locus; $H_o$: observed heterozygosity; $H_e$: expected heterozygosity; $F_{st}$: intrapopulation fixation index. * Mountain agroecosystems (= Rif and North west zones); Oasis agroecosystems (= oases of the South Morocco).
zone appears to be slightly atypical, a feature which could have been predicted. Indeed, the region is the most affected by neighboring cities and as such represents a less traditional agroecosystem, slightly blurring the picture.

The pattern of isolation by distance, with no clines in diversity, is a signature of a genetic equilibrium situation, with no trace of a past colonization process. This feature and the quasi-absence of widespread varieties, is suggestive of a cultivation system based on varieties that originated locally, mainly from the local gene pool.

Table 3 Pairwise $F_{ST}$ values between samples from the different geographic zones

|        | North west | Rif | North center | Center | Moulouya |
|--------|------------|-----|--------------|--------|----------|
| Rif    | 0.028**    |     |              |        |          |
| North center | 0.026* | 0.046** |          |        |          |
| Center | 0.021***   | 0.030*** | 0.025*    |        |          |
| Moulouya | 0.027***  | 0.031** | 0.026*    | 0.017** |          |
| South  | 0.038***   | 0.068*** | 0.042*    | 0.018*  | 0.029**  |

*p < 5.10^{-5}, ** p < 10^{-6}, *** p < 10^{-9}

Variatel and genotypic diversity in mountain and oasis agroecosystems

Quite interestingly, traditional mountain agroecosystems (North west and Rif zones) presented much more varietal diversity than traditional oasis agroecosystems (South zone) (Table 2, Figure 2B and 2C). However they presented almost identical numbers of alleles. This result was obtained despite our sampling only 21 trees in the oases against 141 trees in the North west and Rif zone. This suggests that fig varietal and genetic diversity available in oases is threatened, maybe due to their small surface, while the one available in the mountain agroecosystems will be more resilient.

Conclusions

Traditional Moroccan agroecosystems contain substantial fig varietal and genetic diversity. While fig varieties are true clones and not landraces [16], the distribution of differences between genotypes shows that this diversity arose through sexual reproduction and only marginally, through somatic mutation. Hence the silver bullet hypothesis of instantaneous domestication of clonal plants [13] does not apply, at least today, to fig. In that

![Figure 4 Separation of genotypes according to zone of origin on the two first axes of the Factorial Correspondence Analysis](image)

The Southern zone (in red) and the Rif zone are separated (in blue).
perspective fig is similar to other clonally propagated plants from other parts of the world for which sexual reproduction has been important and often still is. Such species include for instance Cassava [33] and Agave [34] in America or Enset [35] in Africa. Further, in fig, sexual production of new varieties almost obligatorily involves crosses with wild figs. Indeed, it is a dioecious species, and male figs used for pollination are collected on any tree in the neighborhood, and when male figs are cultivated within a village, their potential genetic qualities for siring agronomically interesting crops is not taken into account. Preliminary data from the Rif zone confirms close genetic relationship between local varieties and wild growing fig trees. As such fig cultivation in its native range fits the global picture of frequent hybridization of cultivated plants with their wild relative [36].

However the case of fig is particular as new varieties must (almost) systematically result in the incorporation of hybrids between wild and cultivated plants. We may thus suspect that in all traditional Mediterranean agroecosystems located within fig natural habitat, cultivated figs and wild growing figs locally form a single evolutionary unit. Hence such traditional agroecosystems are effectively incubators of fig variety diversity in a dynamic incorporating wild growing as well as cultivated trees. This is not always the case in clonally propagated plants. For instance, while sexual reproduction seems to be most important in traditional Cassava cultivation, genetics allow to trace its origin to a single region of the range of its progenitor, Manihot flabellatus [37]. The domestication process of monoecious and dioecious plants may turn out to be quite different.

In a context of ongoing rapid climatic change, the nutritional quality, and toxicity of crops may change dramatically [38]. A dynamic management of genetic resources as observed here in traditional agroecosystems may prove essential for responding to such new challenges.

**Methods**

**Fig sampling**

Traditional agroecosystems are still present in Morocco, in the Rif and Atlas mountains in Northern and central areas and in oases in the South east. A survey in the Rif agroecosystems showed that 28 crop species were cultivated including 14 fruit species [31]. A high diversity of fruit crops was also observed in the South Moroccan oases.

Field trips to collect plant material covered all territories of Morocco presenting traditional agroecosystems (Figure 1). They were done in June and August-September in order to observe first or second crop figs, respectively (fig varieties produce either both first crop and second crop or only the second crop). Collections were made in 2005 and 2006. This allowed characterizing the different varieties and establishing their geographical range. Field observations and some genetic data (Achtak et al. unpublished) had shown that within the range of

![Figure 5 Pairwise kinship coefficient between genotypes as a function of geographic distance](http://www.biomedcentral.com/1471-2229/10/28)
each prospection site or village, each variety corresponded generally to a single genetic clone. The sampling strategy could therefore be focused on diversity, using pomological observation following the IPGRI recommendation [39] and interviews with farmers. Thus, for each prospection site, we sampled one individual of each of the cultivated varieties. When we had a doubt on the perfect identity of vegetative and pomological traits within a variety within a site, or when a farmer suggested that there were two types within a variety, then we collected both forms. Hence genetic homogeneity within variety was assessed within site when there was any hint of a doubt, and systematically, among sites. Local variety names were noted as given by farmers; photographs and GPS coordinates were recorded as references for each collected fig tree (see Additional File 1). The photographs allowed confronting a posteriori genotypic identity with morphological similarity. Six major geographical zones were surveyed (North west, Rif, North center, Center, Moulouya valley and South; Figure 1) and 277 trees representing 119 denominations were sampled.

DNA extraction and SSR genotyping
Total genomic DNA was extracted from 200 mg of fresh young leaves of the 277 sampled fig trees using the DNeasy Plant Mini Kit (QIAGEN) according to the supplier’s instructions with the following modification: 1% of Polyvinylpyrrolidone (PVP 40,000) was added to the buffer AP1.

We selected 17 loci among the developed SSR markers [22-25] based on their polymorphism and ease of scoring following the screening of 16 distinct Mediterranean varieties.

Microsatellite amplifications were performed according to the protocol described by Khadari et al. [40]. SSR genotyping was conducted in an automated capillary sequencer (ABI prism 3130XL). Analyses were performed using the GENEMAPPER V3.7 software.

Data analysis
For each SSR locus, alleles were detected and identified by locus name and allele size in bp. Genetic distances between fig genotypes were estimated according to the Jaccard similarity coefficient and UPGMA algorithm using a program developed by J. Brzustowski http://www.biology.ualberta.ca/jbrzusto/cluster.php. The corresponding phenogram was drawn based on the software Treeview 6.1. Discriminating power, D, was calculated for each SSR locus as $D_j = \sum p_i [(Np_i-1)/(N-1)]$ [41] where $p_i$ was the frequency of the $i$-th molecular pattern revealed by locus $j$, and $N$ was the number of genotypes. We used the $D_j$ values to compute the exact probabilities of getting at least one pair of genotypes differing only at 0, 1, 2, 3, 4, 5 and 6 loci.

The number of alleles per locus ($A$), observed heterozygosity ($H_o$), expected heterozygosity ($H_e$) and Wright’s fixation index ($F = 1 - H_o/H_e$) were computed using the software Genetix 4.5 [42]. Genetic diversity was compared among geographic zones using parameters corrected for sample size [43]. Genetic differentiation between populations was assessed using $F_{ST}$ values and the software Genepop 3.1 [44]. The significance of population differentiation was estimated using exact tests [45].

To assess genetic isolation by distance, spatial genetic structure was investigated using a spatial autocorrelation method. Genetic relationships between all pairs of genotypes were regressed on the linear and the logarithmic geographical distance using the software SPAGeDi [46]. The kinship coefficient of Loiselle et al. [47], robust against the presence of low frequency alleles, was used. Significance of the regression coefficients was assessed through 10,000 permutations.

List of Abbreviations
BP: Before Present; pers. comm.: personal communication; DNA: Deoxyribonucleic acid; FCA: Factor Correspondence Analysis; GPS: Global Positioning System; IPGRI: International Plant Genetic Resources Institute; pb: base pair; PCR: Polymerase Chain Reaction; PVP: Polyvinylpyrrolidone; SSR: Simple Sequence Repeat; UPGMA: Unweighted Pair Group Method with Arithmetic mean.

Additional file 1: List of the studied fig trees. This table provides the list of studied fig trees with indications on their sampled geographic zone, sub-zone, site, name, SSR profile and the GPS coordinates.

Click here for file [http://www.biomedcentral.com/content/supplementary/1471-2229-10-28-S1.XLS]

Additional file 2: List of groups of closely related genotypes with skin color fruit. This file describes a list of groups of closely related genotypes differed only by 1 to 3 alleles and considered to be somatic variants of a single clone.

Click here for file [http://www.biomedcentral.com/content/supplementary/1471-2229-10-28-S2.DOC]

Additional file 3: Cases of synonymy. This file describes the cases of synonymy (several variety names for one genotype) observed among cultivated fig trees in Morocco.

Click here for file [http://www.biomedcentral.com/content/supplementary/1471-2229-10-28-S3.DOC]

Additional file 4: Cases of homonymy. This file describes the cases of homonymy (several genotype names for one variety name) observed among cultivated fig trees in Morocco.

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Authors’ contributions

BK designed and coordinated the study. HA, MA, AO and BK performed fig sampling. HA, SS and BK carried out the molecular analysis. HA, FK and BK performed the statistical analysis and wrote the first draft of the manuscript. All authors participated in the draft finalization and approved the final manuscript.

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