



Figure 2: GBIF Data Portal, showing the plant genetic resources data network (<http://data.gbif.org/datasets/network/2/>, visited 10 July 2012). Currently the passport data for more than 2 million genebank accessions (2,180,554) are made available by genebanks through the GBIF distributed data infrastructure.

with research data sets published by the project partners using the BioCASE data publishing toolkit.

GCP Crop Ontology

The International Agricultural Research Centers (IARC) of the CGIAR have developed a Crop Ontology (CO) designed to provide a controlled vocabulary set for some of the economically important crops (Shrestha *et al.*, 2010). The Crop Ontology is based on the Gene Ontology (GO) (Ashburner *et al.*, 2000), Plant Ontology (PO) (Jaiswal *et al.*, 2005), MIAME-Plant (Zimmermann *et al.*, 2006), and the Multi-Crop Passport Descriptors (FAO/IPGRI, 2001).

GENEBANKS IN THE CONTEXT OF BIODIVERSITY INFORMATION

The First Genebanks to Join GBIF

The Global Biodiversity Information Facility (GBIF) was established in 2001 (GBIF, 2001) as an inter-governmental initiative to facilitate free and open access to biodiversity data online. During 2004, the Nordic Gene Bank (NGB, reorganized in 2008 as the Nordic Genetic Resource Center, NordGen), the Polish genebank in Radzików, and the German genebank in Gatersleben were the first genebanks to join the GBIF network (Knüpffer *et al.*, 2004). GBIF is a distributed biodiversity information network based in part on the information standards defined by the Biodiversity Information Standards organization (TDWG; Taxonomic Databases Working Group, 1985–2006). At the time when the first genebanks joined GBIF, TDWG were developing two alternative biodiversity collection data exchange formats, both potentially

suitable for genebank accessions. These were the Darwin Core (DwC, version 2) and the Access to Biological Collections Data (ABCD version 1.20) (Berendsohn, 2005). During 2005 and in collaboration with the ABCD task group, the standard genebank descriptors (MCPD) were mapped to the corresponding ABCD terms, or added as new descriptors to an updated version of the ABCD (version 2.06) (Berendsohn and Knüpffer, 2006). This new development of data interoperability between the crop data sets and the TDWG data-sharing standards opened the possibility for utilization of the GBIF data infrastructure by the PGR community for its own interoperability tasks. It was now possible to start the implementation of the data-sharing toolkits from the GBIF and the TDWG communities in the EURISCO network for Europe (Endresen *et al.*, 2006) (Figure 2). The BioCASE (Berendsohn, 2002) data publishing toolkit was installed at 15 genebanks located in different parts of the world and a demo data portal²³ was developed in 2006 to interact with these distributed web services established by the BioCASE installations.

Dublin Core Metadata Initiative (DCMI)

The DCMI was initiated at a joint workshop between the Online Computer Library Center (OCLC) and the National Center for Supercomputing Applications (NCSA) on metadata semantics held in Dublin (Ohio, USA) in March 1995. The output from this workshop was called "Dublin Core metadata" based on the location of the workshop. The original target was to develop a

²³ <http://www.nordgen.org/portal/index.php?scope=chm>.

small, common set of metadata elements to describe Web content. The original Dublin Core elements (or terms) were: *Subject, Title, Author, Publisher, OtherAgent, Date, ObjectType, Form, Identifier, Relation, Source, Language, and Coverage*. The Dublin Core was designed to be extensible (Weibel *et al.*, 1995).

Darwin Core

Natural history museums in the USA started early to develop information networks with distributed query systems using the Internet. The Species Analyst project was initiated in 1997 and coordinated from Kansas University (Vieglais *et al.*, 1998; Peterson *et al.*, 2003). The first version of the Darwin Core list of terms was developed in 1999 by the Species Analyst project (Stein and Wieczorek, 2004). The Mammal Networked Information System (MaNIS) was initiated in 1999 and established in June 2002 a distributed information network between 17 North American mammal natural history collections. The MaNIS network was developed in parallel with the Distributed Generic Information Retrieval (DiGIR) data publishing toolkit and contributed to the development of the next version of the Darwin Core (DwC version 1.21) (Stein and Wieczorek, 2004). The current version of the Darwin Core²⁴ is more different from the earlier versions than the previous versions are from each other, and was ratified and published by TDWG in October 2009. Darwin Core can be seen as an extension to the standard Dublin Core metadata terms and is designed to be extensible. Darwin Core provides stable semantic definitions of terms for sharing information on biological diversity (Darwin Core Task Group, 2009a; Wieczorek *et al.*, 2012).

Darwin Core Archive (DwC-A)

The GBIF IPT Task Force introduced the Darwin Core Archive format (Döring *et al.*, 2011). The DwC-A is based on the Darwin Core text guidelines²⁵ with core entities linked in a one-to-many relationship to records in extensions. The core entities and the entities in each of the extensions are presented as *fielded text* such as comma separated values (CSV) or tab delimited values (TAB), with one file for the core and one file for each extension. Each record in an extension file is linked to one of the records in the core file. A metafile (meta.xml) describes the structure of the DwC-A including a mapping of the columns in the core and extension files to terms declared by published vocabularies. The DwC-A is created as a zip archive including the

metafile, core and extension files. A resource metadata document (by default named EML.xml) can be included in the zipped archive or referenced with a link to an online metadata resource describing the dataset. GBIF recommends providing metadata using the *GBIF Metadata Profile*²⁶ (Ó Tuama *et al.*, 2011), which is based on the EML²⁷ (ecological metadata language) (Michener *et al.*, 1997; Fegeaus *et al.*, 2005). Sharing the entire dataset in this manner as a DwC-A allows for simpler and more efficient data transfer compared to web service interfaces provided by toolkits such as the BioCASE and DiGIR (Darwin Core task group, 2009b; Döring *et al.*, 2011).

GBIF Knowledge Organization System (KOS)

Recently GBIF has convened expert task groups for providing recommendations on implementing knowledge organization systems (KOS) (Catapano *et al.*, 2011; Lapp *et al.*, 2011), metadata standards (Jones *et al.*, 2009), and persistent identifiers (Cryer *et al.*, 2010; Richards *et al.*, 2011) for biodiversity information resources. One of the recommendations from these task group reports was the implementation of persistent identifiers for each individual vocabulary term concept. When no previously established persistent identifier was available for a term, they recommended a new persistent identifier to be issued. An emphasis was also made on reusing existing terms and concepts wherever possible. These recent guideline principles align very well with the approach that was followed when the Darwin Core extension for genebanks was developed.

[F]lat vocabularies should be developed so that they are reusable as a terminological foundation for semantically richer vocabularies or ontologies (Catapano et al., 2011:3).

The World Wide Web Consortium (W3C) chartered a task force in 2004 for the development of guidelines for managing Resource Description Framework (RDF) vocabularies of terms (W3C, 2006). The task force recommended the following general principles: “(1) use URIs for naming; (2) provide readable documentation; (3) articulate maintenance policies; (4) identify versions; and (5) publish a formal schema” (Kendall *et al.*, 2008). The W3C publishes two dedicated vocabularies with terms for the description of such vocabularies, namely the RDF vocabulary description language

²⁴ <http://www.tdwg.org/standards/450/>.

²⁵ <http://rs.tdwg.org/dwc/terms/guides/text/index.htm>.

²⁶ <http://rs.gbif.org/schema/eml-gbif-profile/1.0.1/>.

²⁷ <http://knb.ecoinformatics.org/software/eml/eml-2.1.0/index.html>.

(RDFS; Brickley *et al.*, 2004) and the simple knowledge organization system (SKOS; Miles and Bechhofer, 2009). Terms from the germplasm vocabulary are described using a combination of properties from both RDFS and SKOS.

RESULTS

During 2008, work was started at GBIF for a major upgrade of the data publishing toolkit for the GBIF network. The new tool was named GBIF Integrated Publishing Toolkit (IPT) and is based on the Darwin Core (DwC). The Darwin Core was at the time under revision by TDWG for a new version scheduled to be ready in 2009. While the ABCD standard is very comprehensive with several thousand terms, the Darwin Core standard implements a more limited set of core terms with domain-specific terms organized in a number of published extensions. There was, as of 2008, no Darwin Core extension to ensure full interoperability with the genebank information requirements. During a Darwin Core workshop in Copenhagen (hosted by GBIF) in January 2009, work was initiated to develop an extension for germplasm to the new revised Darwin Core standard. The Darwin Core extension for genebanks (DwC-germplasm) is required for the rational use of the GBIF IPT in the genebank community (Endresen *et al.*, 2009). The DwC-germplasm provides a comparable piece in the interoperability puzzle as the ABCD version 2.06 provided in 2005 (Berendsohn, 2005; Berendsohn and Knüpffer, 2006) to enable the rational use of the BioCASE toolkit in the genebank community (Endresen *et al.*, 2006).

Darwin Core Extension for Genebanks (DwC-germplasm)

The first draft version of the DwC-germplasm²⁸ was published for discussion at the EPGRIS3 wiki²⁹. The EPGRIS3 (Establishment of a European Plant Genetic Resources Information Infra-Structure, phase 3)³⁰ is an initiative of the ECPGR Documentation and Information Network. The initial development of the DwC-germplasm at the EPGRIS3 wiki attracted feedback and suggestions from the ENSCONET (European Native Seed Conservation Network) project regarding additional terms for *in situ* conservation of genetic resources. The Millennium Seed Bank proposed additional terms to describe *ex situ* germplasm conservation management routines. After receiving further feedback from some other communities outside the

European genebank community, a new DwC-germplasm project home page³¹ was established at Google Code. The Google Code site replaced the EPGRIS3 Wiki as the official home page for the DwC-germplasm vocabulary of terms. Future modifications and eventual additional terms to the DwC-germplasm will be discussed and agreed here before they will be passed on to be consolidated within the genebank community and eventually included in the official version of the extension. The official normative version of the Darwin Core extension for genebanks is published and maintained at "<http://purl.org/germplasm/germplasmTerm.rdf>".

The terms included in the DwC-germplasm vocabulary are organized into 11 groups declared by the SKOS/RDF resource as "skos:Collection" resources. These groups can be seen as entity types and are declared as "rdfs:Class" resources. The germplasm terms describe properties with an intentional loose connection to these entities. The terms organized under *dataset* (1), *taxon* (2) and *collecting event* (3) were imported from Darwin Core and Dublin Core with no new terms declared by the germplasm extension. The description of the *specimens* (genebank accessions) (4) maintained in living collections was supplemented with terms related to the storage conditions and the "biological status of sample" defining the cultivation status ranging from wild plants via landraces and primitive crops to the modern so-called advanced cultivars. The terms for description of the *breeding or domestication event* (5) for crops are unique for the germplasm extension with no overlapping terms imported from the Darwin Core. The same orthogonality of terms in relation to Darwin Core, applies for terms organized to the *acquisition event* (6) describing the donation and sharing of living germplasm material between genebank collections, and also for the *safety duplication event* (7) describing the backup storage of germplasm material at multiple locations. The terms for describing *international treaties and regulations* (8) governing the access and ownership of germplasm material are also unique to the germplasm extension. The Darwin Core terms organized as "*MeasurementOrFact*" (9) were also supplemented and reorganized by the germplasm extension. The germplasm extension provides terms for linking to external resources describing the *measurement method* (trait descriptor) (10). External trait descriptions include measurement methods described and declared by ontologies such as the crop ontology (CO) (Shrestha *et al.*, 2010), plant ontology (PO)³² (Jaiswal *et al.*,

²⁸ <http://rs.nordgen.org/dwc/>.

²⁹ <http://www.nordgen.org/epgris3/wiki/index.php/DwC.Germplasm>.

³⁰ <http://www.epgris3.eu/>.

³¹ <http://code.google.com/p/darwincore-germplasm/>.

³² <http://www.plantontology.org/>.

2005), the phenotypic quality ontology (PATO)³³ and the plant trait ontology (TO)³⁴ (Jaiswal *et al.*, 2002). Some terms were also added related to the description of the *measurement experiment* (11) including the time and location of the experiment. The goal was the maximum reuse of existing terms from existing vocabularies and ontologies and the minting of new terms only when no comparable terms were found.

Darwin Core Archive (DwC-A) Extension for Genebanks

For the terms declared as part of the DwC-germplasm to be made available to the GBIF infrastructure software tools such as the GBIF IPT and to be included into data sets shared using the DwC-A data-publishing format (Döring *et al.*, 2011), the relevant XML application lists were developed. The GBIF Resources Registry provides the formal specifications for these XML applications in the format of a XML schema³⁵. The GBIF Vocabulary Server³⁶ provided a software tool to assist in the development of the DwC-A extensions³⁷ for the DwC-germplasm terms. The final XML applications required for including the germplasm terms to DwC-A dataset resources were published at the GBIF Resources Registry³⁸.

Deployment of DwC-germplasm in GBIF IPT

In 2010, GBIF, NordGen and Bioversity International initiated a feasibility study to evaluate how the GBIF infrastructure can meet the needs of the European genebank community (Gaiji *et al.*, 2010). This feasibility study was also coordinated with the ECPGR Documentation and Information Network for the genebank community in Europe (Maggioni, 2010). The prototype GBIF Integrated Publishing Toolkit (IPT version 1.0) was installed in five genebanks within the European plant genetic resources catalogue (EURISCO) using the DwC-germplasm extension. These were the national genebanks in the Russian Federation (Vavilov Institute, Sankt Petersburg), Germany (IPK Gatersleben), Czech Republic (Crop Research Institute, Prague), the Netherlands (Wageningen University and Research Center, Centre for Genetic Resources), and within the Nordic and Baltic countries (genebank database hosted from the Nordic Genetic Resources Center).

While the prototype version of the IPT software caused some problems of instability during installation (1), the mapping of the genebank data sets to Darwin Core, including the gene bank extension (2) and finally the registration to the GBIF GBRDS (3) was completed satisfactorily. The hardware requirements and in particular the demands for internal memory were a major barrier encountered during most of the installations. The experiences from the genebank feasibility study provided the IPT development team at GBIF with feedback and suggested improvements leading from the different prototype versions to the new version 2. The hardware requirements for the new version of the IPT software (version 2) have been significantly reduced and solve all of the issues encountered and reported from the genebank feasibility project. The project resulted in a positive evaluation and the genebank community has started the initial plans for a second feasibility study to evaluate the IPT version 2 at other genebanks in Europe.

Updates to the Vocabulary of Germplasm Terms

Following the experiences from the first draft version of the DwC-germplasm (version 0.1) and the IPT feasibility study at some of the European genebanks, some updates to the germplasm terms were made. Based in part on the recommendations from the GBIF KOS task group (Catapano *et al.*, 2011; Lapp *et al.*, 2011) and the initial work by the Vocabulary Management Task Group³⁹ (Endresen *et al.*, 2012a), the DwC-germplasm terms were declared as a RDF/SKOS vocabulary. The recommendations from the World Wide Web Consortium on best practices for management of RDF vocabularies (Kendall *et al.*, 2008) provided other useful principles for the upgrade of the germplasm terms.

The updated vocabulary of germplasm terms⁴⁰ was moved to the PURL (persistent uniform resource locators) namespace to improve the commitment of long-term persistence. PURL is managed by the Online Computer Library Center (OCLC) to provide persistent and resolvable identifiers for online resources.

RDF Vocabulary Maintenance Policy

The DwC-germplasm vocabulary of terms is extensible and new terms can be added to the DwC-germplasm namespace in the future. Terms can evolve through refinement in response to deployment and testing. Refinements to the definition and description of a term will as far as

³³ http://obofoundry.org/wiki/index.php/PATO:Main_Page.

³⁴ http://www.gramene.org/plant_ontology/.

³⁵ <http://rs.gbif.org/schema/extension.xsd>.

³⁶ <http://vocabularies.gbif.org/>.

³⁷ <http://vocabularies.gbif.org/extensions/>.

³⁸ <http://rs.gbif.org/extension/nordgen/>.

³⁹ <http://community.gbif.org/pg/groups/21382/vocabulary-management/>.

⁴⁰ <http://purl.org/germplasm/germplasmTerm.rdf>.

possible maintain the semantic meaning of the term. In situations when the original semantic meaning of a term is jeopardized, the term will be deprecated and replaced by a new term. The previous versions of terms will be maintained in a separate vocabulary describing the history of the terms. All terms are described by a label (skos:prefLabel), a definition (skos:definition), some examples (skos:example) and a note (skos:note) explaining the scope and how to use the term. These natural language descriptions are only available in English. A separate vocabulary might be developed to provide descriptions of the terms expressed in other natural languages than English.

promotes the interoperability of biodiversity data with information from other domains. The Darwin Core set of core terms includes some terms from the Dublin Core ‘terminology’⁴². But more important than the shared terms, is the shared framework to describe and implement the terms in applied solutions. The same principles apply to the benefits of building a genebank extension based on the Darwin Core, or adapting other solutions from outside the germplasm community network. By following a few ways and guidelines for ‘best practices’, the genebanks can with few efforts adapt tools and principles developed in other communities for efficient use in their own information network (Knüpfer *et al.*, 2007).

DISCUSSION

The Darwin Core standard is itself an extension of another standard, the Dublin Core Metadata Initiative (DCMI)⁴¹. The Dublin Core provides a bridge to ensure low-level interoperability between wide ranges of metadata standards. Implementing Darwin Core as an extension to the Dublin Core

The achieved compatibility of data standards between PGR and biodiversity collections allows integrating the worldwide germplasm collections into biodiversity information networks. Using GBIF technology (and contributing to its development), the PGR community can easily



Figure 3: During the 2010 feasibility study for the European genebank community, the prototype GBIF Integrated Publishing Toolkit (IPT) was installed at the national genebanks in the Russian Federation, Germany, Czech Republic, the Netherlands, and at the Nordic Genetic Resource Center (hosting the genebank database for the Nordic and Baltic countries).

establish specific PGR information networks without creating its own technology (Knüpffer et al., 2007:7).

By finding a few common ways and guidelines for ‘best practices,’ genebanks can with fewer efforts adapt tools and principles developed in other parts of their ‘own’ community for efficient use across the entire germplasm information network.

Automatic Data Exchange Mechanisms

Many of the present data exchange mechanisms in use in the genebank community rely on laborious and repeated transformations of the original genebank data sets into the agreed standard formats. The genebanks in Europe regularly produce an updated subset from their information system complying with the EURISCO data exchange format (based on the Multi-Crop Passport Descriptors). Then the subset from each genebank in a country is combined into a so-called National Inventory and uploaded to the central EURISCO data portal (hosted by Bioversity International). The European Central Crop Databases (ECCDB) also request from each genebank to extract a similar subset from their information system.

Many ECCDBs ask for data on selected descriptors from the Bioversity Crop Descriptor lists in addition to the MCPD. The ECCDBs are limited in scope each to a different crop species and ask thus for a different set of additional crop-specific descriptors. While the EURISCO has implemented an online data upload tool to receive the updated national inventories, the updated subsets for the ECCDBs are often exchanged as email attachments. The CGIAR genebanks share similar subsets from their information systems with the System-wide Information Network for Genetic Resources (SINGER)⁴³. The FAO WIEWS (World Information and Early Warning System on PGRFA)⁴⁴ also requests, on a regular basis, updated subsets from all genebanks worldwide. The requested format for these subsets is also roughly based on the MCPD standard. The record level data unit is however different, as WIEWS request metadata on stratified groups of genebank accessions, rather than the accession level data requested by EURISCO, ECCDBs and SINGER.

New data exchange mechanisms using web services have the potential to make all these aforementioned data exchange operations fully automatic. And with the new data provider toolkit software packages provided as an open source public

good from the Global Biodiversity Information Facility (GBIF), the required efforts to establish and maintain such fully automatic multi-purpose data flow pathways with web services are getting less demanding and becoming more low-tech to implement. The GBIF Integrated Publishing Toolkit (IPT) is the latest and most user-friendly software package for sharing biodiversity data sets (such as the genebank data sets). The Darwin Core extension for genebanks (DwC-germplasm) provides a necessary ‘plug-in’ to make the new GBIF IPT available for rational use in the genebank community. The genebank community was one of the first biodiversity information networks to develop this type of plug-in to start using the GBIF IPT. It is expected that the experiences from the development and implementation of the DwC-germplasm for the genebanks can provide some examples for other biodiversity information networks to study. With the following section, we aim to describe the steps to follow in order to develop a similar Darwin Core extension in other biodiversity information networks.

HOW TO Create a New Darwin Core Extension

The development and implementation of the Darwin Core extension for germplasm can be used as an example for other biodiversity information communities to develop their own DwC extensions. The following steps have to be carried out:

- (1) The community needs to compile a consolidated list of terms to describe their data domain.
- (2) After finding agreement on the terms with the relevant stakeholders inside the relevant community, these terms should be harmonized and mapped to the standard Darwin Core terms⁴⁵. New terms should only be defined for an extension if they are not already included in the standard core terms. Some of the descriptor terms implemented in a community may be similar to one of the core terms, but with a different formatting or a slightly different semantic meaning. Whenever possible, it is recommended to try to convert the data content for a community descriptor term to follow the definition of one of the standard DwC terms. If a new community term is defined that could have been converted to one of the existing DwC terms, interoperability with biodiversity data sets from other communities will be broken.
- (3) We recommended declaring new terms for your Darwin Core extension using the simple knowledge organization system (SKOS) and

⁴³ <http://singer.cgiar.org/>.

⁴⁴ <http://apps3.fao.org/wiews/wiews.jsp>.

⁴⁵ <http://rs.tdwg.org/dwc/terms/index.htm>.

the resource description framework (RDF/RDFS) vocabulary.

- (4) Darwin Core extensions and other community terminology vocabularies can be published at the GBIF Resources Registry⁴⁶.
- (5) The next step is to create appropriate XML lists including your terms following the GBIF XML schema specifications for Darwin Core Archive extensions. When designing new DwC-A extensions you may mix and match terms from many different term vocabularies. The GBIF Vocabulary Server (Harman *et al.*, 2009) provides a software tool to assist you with defining a correctly formatted DwC-A extension.
- (6) The final DwC-A extension must be loaded to the GBIF Resources Registry before it is available to software tools such as the GBIF Integrated Publishing Toolkit (IPT). You will (at least for now) need to contact the GBIF helpdesk (helpdesk@gbif.org) for assistance with loading your resources to the GBIF Resources Registry.

Please note that after publishing the DwC-A extension to the GBIF Resources Registry, any modifications to the extension (however minor) need to be released with a new version number. The recommendation for the release of new DwC-A extensions based on the DwC-germplasm terms is to include a postfix with the release date (e.g., “germplasm_20120710.xml”).

New terms and concepts should be developed in a collaborative manner allowing for feedback from your community. The Vocabulary Management Group (VoMaG)⁴⁷ currently performs an evaluation of various software tools⁴⁸ to support the collaborative development of new terms including the Semantic MediaWiki⁴⁹, ISocat⁵⁰ and Drupal-based⁵¹ tools. The ratified version for each vocabulary of terms for the description of biodiversity information resources can be registered and deposited at the GBIF Resources Registry. Darwin Core Archive extensions and controlled value vocabularies should be designed to re-use terms from one of the ratified and published flat vocabularies (RDF/SKOS) or from a published ontology (OWL). We also recommend here as a best practice guideline to re-use terms declared by a flat vocabulary when developing new biodiversity ontologies (OWL resources).

Because the genebank community already had established information standards, the development of a draft extension to the Darwin Core (DwC-germplasm), and the subsequent testing of the new prototype information publishing toolkit from GBIF (GBIF IPT), progressed quickly and with relatively few problems.

Evaluation of the Updated GBIF IPT Version 2

The experiences so far from testing the updated version 2 of the IPT are mostly positive. All the graphical features for visualization of the data sets were removed in this version. This simplification was one of the recommendations reported by the genebank feasibility study. This focus of the IPT on data publishing rather than visualization has resulted in substantially improved performance of the toolkit. The new version 2 has also removed the embedded internal database. As a result of this modification, the web services providing various query interfaces to the data sets shared by IPT have also been removed. In particular the web service interfaces providing access with the TAPIR protocol (TDWG access protocol for information retrieval)⁵² (TDWG, 2010), and the OGC (open geospatial consortium) WFS (web feature service) were interesting APIs (application programming interfaces) to the underlying data. However, the simplicity of the IPT with a dedicated design for providing the Darwin Core Archive (DwC-A) data-sharing format makes the IPT a lightweight and efficient software application. There are also other similar data publishing toolkits such as the TapirLink⁵³ and BioCASE⁵⁴ that provide GBIF-compatible services for publishing biodiversity data sets.

Efficient Access to Distributed Germplasm Data Sets Stimulates Novel Uses

Research integrating genebank passport data (georeferenced occurrence data for the original collecting site) with phenotypic measurements (characterization and evaluation data) and with ecological layers has opened new possibilities for a rational utilization of genebank materials (Bhullar *et al.*, 2009; Endresen, 2010; Endresen *et al.*, 2011; Bari *et al.*, 2011; Endresen *et al.*, 2012b) using the *Focused Identification of Germplasm Strategy* (FIGS) approach (Mackay and Street, 2004). The efficient application of the FIGS approach depends on the availability of germplasm passport and trait evaluation data. The analysis of gaps in the

⁴⁶ <http://rs.gbif.org/terms/>.

⁴⁷ <http://community.gbif.org/pg/groups/21382/vocabulary-management/>.

⁴⁸ <http://kos.gbif.org/>.

⁴⁹ <http://semantic-mediawiki.org/>.

⁵⁰ <http://www.isocat.org/>.

⁵¹ <http://drupal.org/>.

⁵² <http://www.tdwg.org/activities/tapir/>.

⁵³ <http://sourceforge.net/projects/digir/files/TapirLink/>.

⁵⁴ http://www.biocase.org/products/provider_software/.

genebank collections to guide the planning of rational germplasm collection expeditions to complement the genebank collections with novel and insufficiently sampled genetic diversity is also dependent on the availability of genebank passport data (Jarvis *et al.*, 2003; 2005; 2009; Ramírez-Villegas *et al.*, 2010). Such data analysis experiments to identify the ecological environment linked to a target trait property, or the genetic gaps of the genebank collections, will of course benefit from occurrence data on crop wild relatives provided from other communities. The value of external data from outside the genebank community, in such studies, strengthens the argument for the development of common semantic data standards (like the Darwin Core) and standardized data exchange protocols (such as the Darwin Core Archive format). Limited access to genebank accession-level information is a bottleneck to the efficient use of genebank material (FAO, 2010), as well as to the development of novel uses for the associated data. The authors of this manuscript propose the Darwin Core extension for genebanks and its implementation in the GBIF Integrated Publishing Toolkit (IPT) as a contribution for an upgrade of the current data exchange mechanism for genebank data sets.

Future Work

After the first experiences with the deployment of the Darwin Core extension for genebanks, a useful next step will be to seek ratification of the extension as a TDWG standard. The genebank community has long and successful experience with the development and maintenance of descriptor standards, in particular through the work at Bioversity International (Bioversity International, 2007; Gotor *et al.*, 2008). However, as discussed above, one of the major achievements with the DwC-germplasm is the interoperability with other biodiversity information standards and communities outside the genebank community. The ratification of genebank standards like the DwC-germplasm in TDWG will contribute to improved information interoperability.

The first version of the DwC-germplasm included the proposed EPGRIS3 descriptors for evaluation and characterization data. The sharing of trait data sets for germplasm has received renewed attention with the second report on the state of the world's plant genetic resources for food and agriculture (FAO, 2010). These descriptors need further work after the first experiences with the sharing of germplasm trait data sets.

The implementation during the last years of new international regulations for the sharing of benefits for the use of plant genetic resources prescribes the reporting of the distribution of seed samples (defined by the ITPGRFA, Annex 1). If the terms to describe and report these seed distributions are developed and included to the DwC-germplasm, then the GBIF IPT could be used to report seed distributions to the Governing Body of the ITPGRFA.

When germplasm data sets are published, the entities are almost exclusively identified using local identifiers. Often the institutes sharing these data sets are identified using institute codes from the FAO WIEWS (world information and early warning system)⁵⁵. The combination of the WIEWS institute code and the local identifier for entities (such as genebank accessions) are generally sufficient to ensure unique identification. One major concern with this practice is that the institution codes are not designed to be globally unique and persistent identifiers (PIDs). For a description of globally unique and persistent identifiers see Coyle (2006), Campbell (2007) and Richards *et al.* (2011). The required combination of the institute identifier and the local identifiers for distinct identification of entities is another major bottleneck for efficient use of information from the germplasm data sets published online today. The *GBIF data publishing framework task group* recommends the publication of biodiversity data sets as citable “data papers” and that each dataset is identified by a PID for consistent data citation (Moritz *et al.*, 2011). The Dryad data repository provides a similar service where the data sets supporting published peer-review papers can be archived and provided with a DOI (digital object identifier)⁵⁶ for consistent citation and persistent data availability (Greenberg *et al.*, 2009; Michener *et al.*, 2011). The persistent identification of data sets could become a major step towards consistent identification of entities in germplasm collections. However, we recommend that also the entities inside the germplasm data sets will be identified using PIDs. It is our opinion that the full potential of using the DwC-germplasm terms when publishing germplasm data sets will be limited without the use of PIDs for consistent identification of entities inside the data sets such as the genebank accessions. Efforts should further be made for the reuse of existing PIDs for such entities.

The Darwin Core extension for genebanks was developed as a flat vocabulary of basic terms. This vocabulary is declared using RDF/SKOS and

⁵⁵ http://apps3.fao.org/wiews/institute_query.htm?i_l=EN.

⁵⁶ <http://www.doi.org/>.

declare only very limited formal semantics for the germplasm terms. Further declarations of formal semantics using the web ontology language (OWL) would improve the interoperability of the germplasm terms in relation to other ontologies declared using OWL. Ontologies declared using OWL provides formal logic constructs that enable logical inferencing. Machine reasoning is a powerful tool in knowledge integration that can give rise to new, inferred knowledge (Allemang and Hendler, 2008). We recommend here the OWL ontology developed as a complement to the flat list of germplasm terms and not as a replacement for the flat RDF/SKOS vocabulary. We recommend here as a best practice guideline to maintain the terms as a flat (SKOS) vocabulary to maximize the potential for reuse of the terms and that multiple OWL ontologies based on the same terms can efficiently be developed for different purposes. We recommend a flexible governance model allowing for differing ontological views to be expressed while reusing the same terms.

CONCLUSIONS

The Darwin Core germplasm extension provides access to the GBIF bioinformatics infrastructure, including the GBIF Integrated Publishing Toolkit (IPT). Using the GBIF IPT and the Darwin Core germplasm extension, genebanks can now share germplasm data sets with each other. This new data exchange mechanism will make the development of distributed germplasm information networks easier. The DwC germplasm extension also provides a frame for implementing a standardized process of data exchange. Implementation of general biodiversity information standards and toolkits will ensure the interoperability of genebank data sets with other biodiversity data sets.

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