

Processing USPTO Patent Data

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Abstract

We describe a completely automated data process designed to consume weekly releases of patent grants and applications distributed by the United States Patent and Trademark Office (USPTO). The process downloads and unpacks the zipped distribution files, parses the raw data into a SQL database, and performs various disambiguations and statistical calculations on the database.

1 Introduction

Patent data plays an invaluable role in research into economic trends, invention, innovation policy and technology strategy. Since the digitization of patent data starting in 1975, though patent data has been freely available through the United States Patent and Trademark Office, it has been difficult to use. We present a substantial improvement in data quality and accessibility over previous third-party re-releases of US patent data. This will not only facilitate further research on up-to-date patent records, but also increase the reproducibility of previous research results.

2 Processing Workflow

The Fung Institute has developed a robust and fully automated toolchain for processing and

providing high quality patent data intended for research, as illustrated in Figure 1.

As data is downloaded from the USPTO weekly patent releases, it is parsed, cleaned and inserted into a SQL database. From this database, assignee and lawyer disambiguations are performed and the patents are geocoded with a location-based disambiguation. The output data from these processes are combined with the historical data from the Harvard Dataverse Network into a single consolidated database. From this database, an inventor-level disambiguation can be performed, and various applications can take advantage of the completed data.

3 Data Sources

The unified patent dataset is composed of processed data from two separate sources: the Harvard Dataverse Network (DVN) [12] collection of patent data from 1975 through 2010 and the weekly distributions of Google-hosted USPTO records [9][10].

3.1 Harvard DVN

The Harvard DVN patent database consists of data drawn from the National Bureau of Economic Research (NBER), weekly distributions of patent data from the USPTO, and to a small extent, the 1998 Micropatent CD product [13]. The schema of the database can be found in the Appendix.

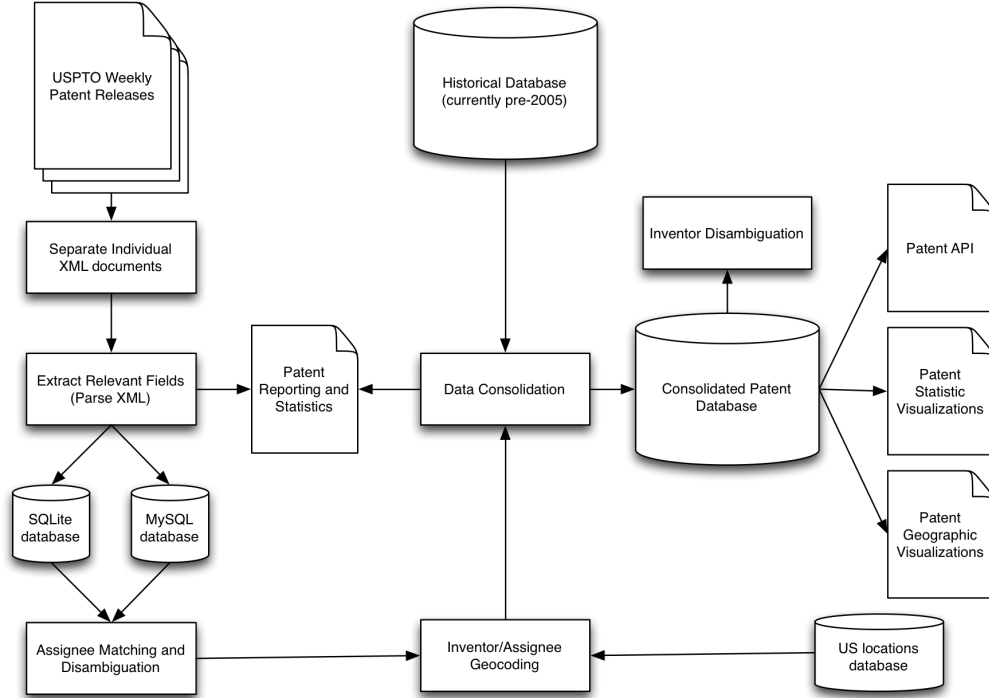


Figure 1: Full patent data process flow

While the Harvard DVN patent database was, prior to the UC Berkeley patent database, the most extensively complete amalgamation of United States patent data, it is not without its problems. Firstly, there is little information as to the actual meanings of the columns in the databases. Without sufficient prior knowledge of patent structure, it is difficult to glean the semantic significance of each column. The names alone are often abbreviated and hard to discern. Secondly, because the DVN database is a combination of several sources into a single database schema, certain patent entries from NBER and Micropatent are incomplete where their data source did not provide all the requisite data. The data obtained from the weekly distributions suffers from being made available in several different formats. The parser that was developed to handle the data is overly complicated and does not handle edge cases well, resulting in missing patent metadata where the parser did not account for a subtle change in format [1]. This is

analyzed in greater detail below.

3.2 USPTO Weekly Distributions

The USPTO distributions take the form of zip archives containing concatenated XML (Extensible Markup Language) documents, each of which contains the full text of each patent grant and patent application issued every week. Prior to 1975, the USPTO used a purely paper-based system before transitioning to a raw-text key-value and later SGML-based key-value store¹. Patent documents were made available in the XML format starting in 2001. Although this data is made freely available, the fact that digital USPTO patent data spans eight different formats and occupies more than 70 GB (compressed) over the 37 years of its existence makes rendering the data into an amenable form a nontrivial problem (see Table 1 and Table 2). Patent application data, though only available in a digital format back

¹Standard Generalized Markup Language

Time Span	Data Format
-1974	paper-based
1975	unknown. Data obtained from Micropatent
1976 - 2001	Green Book (CITE) APS key-value
2001	SGML ST. 32 v2.4
2002 - 2004	Red Book (CITE) XML ST. 32 v2.5
2005	Red Book XML ST. 36 (ICE) v4.0
2006	Red Book XML ST. 36 (ICE) v4.1
2007 - 2012	Red Book XML ST. 36 (ICE) v4.2
2013	Red Book XML ST. 36 (ICE) v4.3
2013 -	Red Book XML v4.4

Table 1: Table of USPTO grant data formats

Time Span	Data Format
-2001	paper-based
2001	XML ST. 32 v1.5
2002 - 2004	Red Book (CITE) XML ST. 32 v1.6
2005	Red Book XML ST. 36 (ICE) v4.0
2006	Red Book XML ST. 36 (ICE) v4.1
2007 - 2012	Red Book XML ST. 36 (ICE) v4.2
2013 -	Red Book XML ST. 36 (ICE) v4.3

Table 2: Table of USPTO grant data formats

to 2001, is nonetheless available in six different formats [14] [15].

4 Parsing

The process of converting the public patent data into a usable form begins with parsing, the manipulation of a document’s grammar and anatomy to extract structured and labeled data. The Fung Institute parser takes as input the weekly USPTO patent distributions and outputs the relevant data into a SQL database. To simplify the problem of parsing the diversity of formats of digital patent data, the current parser addresses only the XML-based documents. At time of writing, the Fung Institute parser is capable of handling patent grants of formats XML v4.0, v4.1, v4.2, v4.3 and v4.4 (spanning 2005 through 2013) and patent applications of formats XML v1.5, v1.6, v4.0, v4.1, v4.2 and v4.3 (span-

ning 2001 through 2013). Grant data prior to 2005 is drawn from a truncated version of the Harvard DVN database.

The code is written in Python 2 [7] and is available on Github [2].

4.1 XML Overview

XML, or Extensible Markup Language, defines a set of rules for encoding documents that seek to facilitate comprehension by both machines and humans. Since the publishing XML 1.0 standard in 1996, the format gained traction due to the minimal size and flexible structure. In its simplest form, an XML document is a collection of elements, which are each composed of tags and content. Tags, such as `<citation>`, lend semantic structure to a document and allow a reader to determine the significance of the content that follows. An element is a logical component that begins and ends with tags (e.g. `<citation>` and

```

<applicants>
  <applicant sequence="001" app-type="applicant-inventor">
    <addressbook>
      <last-name>Roach</last-name>
      <first-name>Richard</first-name>
      <address>
        <city>Schaumburg</city>
        <state>IL</state>
        <country>US</country>
      </address>
    </addressbook>
  </applicant>
  <applicant sequence="002" app-type="applicant-inventor">
    ...
  </applicant>
</applicants>

```

Figure 2: Sample inventor element from XML v4.2 schema

</citation> and contains either regular text or additional, nested elements. An example of an element can be found in Figure 2.

4.2 Parser Method

The Fung Institute parser adopts a novel approach to the problem of extracting data from XML documents. As XML documents are fed to the parser, they are transformed from XML’s canonical tree-based organization into modified Python dictionaries. Typical XML parsers must make certain assumptions about the nesting and placement of tags and must contain careful allowances for missing, mislabeled, or unexpected tags. The Fung Institute parser circumvents this issue by not requiring a detailed specification of the data to be extracted, instead relying on general descriptors of the location of needed data. This makes the parser more robust and able to handle small schema changes without adjustment, therefore reducing the number of potential runtime errors. The existence of such an engine also expedites the development of additional parsers that handle subsequent changes to the USPTO patent XML schemas.

The XML parsing engine reduces the amount

of explicit error checking code while making the source code concise and easy to understand. The engine is easily configurable, and can be directed to automatically download and parse patents in a given date range, apply arbitrary post parsing steps, and deliver the results to a database.

4.3 Data Idiosyncracies

While the USPTO patent data is public and freely available, it is not without its problems.

There is inconsistent usage of HTML idioms and escaping. Underscores, ampersands, em-dashes and brackets – to name a few – are not expressed as literal characters in the raw XML, and care must be taken to translate sequences such as `&` and `_{—}` so that the extracted data is human-readable.

Accents within names are irregularly represented and follow differing standards. Accented letters are either missing (e.g., “Rémy” becomes “R my”) or replaced by description (e.g. “Rémy” becomes “R acute over e my”) or replaced by the same letter without accent (e.g., “Rémy” becomes “Remy”). All three versions of the name “Rémy” are found across the DVN databases and USPTO weekly publications.

Last name prefixes such as “van der” and titles such as “Esquire” are varyingly included in either the `<first-name>` or `<last-name>` tags, which complicates the parsing of names into a consistent form.

The document numbers of patents are inconsistently prefixed with letters representing the type of document, and are occasionally padded with a leading “0”. These eccentricities exacerbate the logical complexity of the parser, but must be handled in order to maintain consistent notation that enables the reliable tracking of references to documents.

These issues are handled by the Fung Institute parser, and are discussed at length in a previous Fung Institute publication [3].

5 Database

One of the main purposes of the patent processor project is to provide a usable database of relevant patent data. This database should facilitate the retrieval of patent records, citations, inventors, lawyers, assignees, and other patent-related data. The linked nature of these types of records suggests that a relational database model would be most suited to the data, which motivated the decision to model patent data in SQL. SQL, or Structured Query Language, is a language designed for managing data held in a relational database.

Because the majority of the data processing pipeline is written in Python, it is hard to integrate otherwise easy-to-use SQL code. There are multiple flavors of SQL – among them, SQLite and MySQL. SQLite simplifies local development because the whole database is represented as a single efficiently-sized file that can be copied, moved and manipulated much like a traditional file. However, it is hampered by a lack of support for more complex SQL features, and has poor support for concurrent users (e.g. multiple processes attempting to access the same database). MySQL offers advanced SQL features (such as `LEFT OUTER JOIN`) and scales to multiple users

and large amounts of data much easier than SQLite, but requires more specialized knowledge to use and access. MySQL is more suited for production environments, whereas SQLite is better for development. We want to be able to easily switch between these two flavors of SQL depending on our purpose without having to develop multiple branches of database integration.

5.1 SQLAlchemy

SQLAlchemy [6] is a Object Relational Mapper (ORM) for Python that seeks to abstract away the differences between SQLite, MySQL, and other SQL-based relational databases. The SQLAlchemy ORM maps Python classes to an underlying SQL database such that the database can be manipulated as though it were a native Python object. This means that the object model and the database schema can be decoupled, effectively removing the need for separate lines of development for each possible database engine.

Database-related code written using SQLAlchemy is much cleaner and easier to work with than the traditional, kludgy idioms. In the case of SQLite, the normal Python module requires the user to excute strings of SQL code:

```
number = ‘‘%s’’ % patent_number
connection.execute(query)
```

Not only does this require the programmer to know SQL syntax, but this paradigm leaves the database open to SQL injection, wherein unintended and possibly malicious code is executed on the SQL database. For example, here, we are operating on the assumption that the variable `patent_number` contains a valid patent number. It could actually contain the string `’’; delete from Patent;--`, which would terminate the original `select` statement, delete all entries from the Patent table, and then exit as though nothing had happened. To avoid such attacks, it is necessary to sanitize all SQL strings to make sure they contain valid and safe queries.

```

number = “%s” % patent_number
result = connection.execute(query)
patent_id = result[3]
query = 'select * from assignee \
where patent_id = “%s” % patent_id
connection.execute(query)

```

Figure 3: Finding assignees for a patent using traditional Python-SQL

```

filter_by(number = patent_number)
patent.assignees

```

Figure 4: Finding assignees for a patent using SQLAlchemy

SQLAlchemy obviates the need to implement such verbose security methods. The SQLAlchemy equivalent to the above query is:

```
filter_by(number = patent_number)
```

Immediately, we can see that this code is much simpler and cleaner. When SQLAlchemy accepts string input, as with the `patent_number` variable here, it automatically escapes all significant characters like semicolons and apostrophes, essentially nullifying the possibility of SQL injection attacks.

SQLAlchemy further simplifies the handling of foreign keys and complex joins between tables, and can even implement these features over database engines (such as SQLite) that do not normally have them. Consider Figure 3 versus Figure 4.

5.2 Limitations

The nice features of SQLAlchemy come at a price. The higher level interface to the SQL database requires a nontrivial amount of book-keeping. Foreign keys lookups and checks introduce a certain amount of overhead, so when a process loops through a list of database items, multiple SQL queries can be executed against the backend for each object if the process asks for linked objects.

SQLAlchemy offers tools to help reduce the number of individual queries sent to the underlying database, but there is an inescapable overhead to using an ORM over the raw SQL.

5.3 New Schema

We wanted to have a highly-linked database that would make it easy for developers to access related information for a given set of patents. The DVN schema, as described in the Appendix, does not take advantage of foreign key relations, and places much manual burden on the user. This was a primary motivating factor in our design, which is summarized in Figure 5.

5.4 Raw vs Processed

If we examine the new database schema, for each of the `inventor`, `lawyer`, `location`, and `assignee` tables, we can see a “raw” version (e.g. `rawinventor`) and a plain version. The `raw` tables contain the inventor, lawyer, location and assignee records *as they appear in the USPTO files*, which means that the naming inconsistencies and misspellings are preserved. These records are run through disambiguation methods of various degrees of rigor, and the cleaned records are stored in the plain tables. See below for a description of these disambiguation methods.

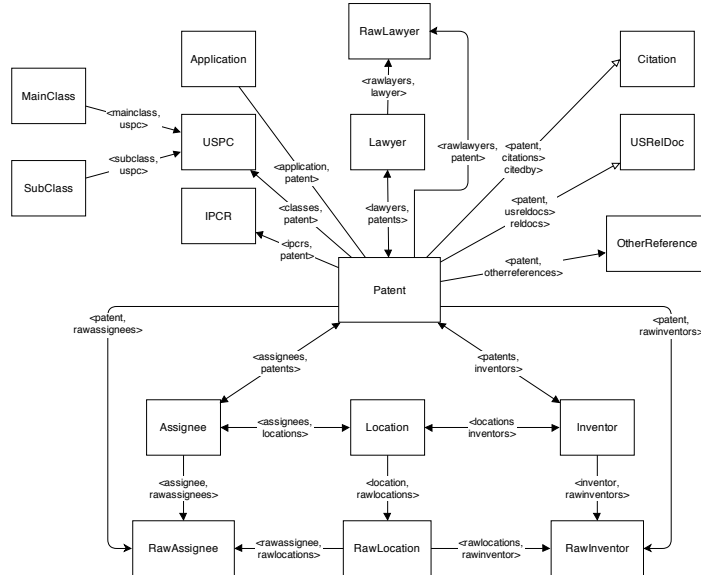


Figure 5: High level view of new database schema

When the cleaned records are inserted, we link them to both the related patent and the raw version using foreign keys in the database, so it is simple to examine groups of related records. See Table 3.

6 Disambiguations

One of the primary problems with conducting meaningful research with USPTO patent data is the high variability in quality. Cities are misspelled or mislisted. Organizations are alternatively abbreviated and listed in full with little modicum of consistency. Inventors, lawyers and assignees will misspell their names, change their names and unpredictably list their middle initials or names. The Berkeley patent database provides facilities to account for these errors, and codifies the disambiguation of such records in order to make possible their accurate retrieval.

6.1 Geocoding

There are over 12 million locations listed in the USPTO patent weekly downloads from 1975 to 2013, with 350,000 unique tuples of

(city, state, country). These tuples follow the typical motif of data problems in the rest of the patent data: incorrect or nonstandard country codes, inconsistent romanization of foreign locations and various misspellings. We resolved the ambiguities in the location data using a proprietary disambiguation technique developed by Google. When new patent data is processed, we run a series of data cleaning processes to correct for some of the common errors, then cross reference with the lookup table [4] obtained through the Google disambiguation.

A detailed analysis of the problems with USPTO location data and our handling of locations can be found through a related Fung Institute publication [5].

Locations are associated with assignees, inventors and lawyers. Typically, a patent record’s “location” is the location of the first inventor listed on the patent.

6.2 Assignees

For a given patent, the assignees are the entities (either organizations or individuals) that have property rights to the patent. The assignee

Table	Access	Value
Patent	<code>patent</code>	US8434162
Inventor	<code>patent.inventors[0]</code>	Thomas H. Stachler
Raw Location	<code>patent.inventors[0].rawlocation</code>	Deyton, OH, US
Clean Location	<code>patent.inventors[0].location</code>	Dayton, OH, US

Table 3: Accessing related raw and clean records. Note the spelling correction in the clean record

records are imperative for firm-level analysis of patent data, and are used for tracking ownership of patents. The weekly releases of patent documents only contain the original assignee of a patent when it was initially granted.

However, it is difficult to obtain accurate results for simple (and necessary) questions such as *“which patents are owned by firm X?”* because of the pandemic inconsistency of spellings. A cursory search for assignee records that resemble General Electric yields the following:

- General Electric Company
- General Electric
- General Electric Co..
- General Electric Capital Corporation
- General Electric Captical Corporation
- General Electric Canada
- General Electrical Company
- General Electro Mechanical Corp
- General Electronic Company
- General Elettric Company

This is not even a complete list of all the (mis)representations of General Electric, but already we can see the potential issues with trying to get accurate results.

We do not yet provide fully featured entity resolution for assignee records, but we do maintain a preliminary disambiguation of the records that corrects for minor misspellings. We do this by applying the Jaro-Winkler [8] string similarity

algorithm to each pair of raw assignee records. Two records that are within a certain bound of similarity are considered the same, and are linked together.

It is not tractable to perform pairwise computation on each of the 5,850,531 raw assignee records in the database (at time of writing), so we group the assignees by their first letter, and then perform the pairwise comparisons within each of these blocks. This allows us to hold a smaller chunk of the assignees in memory at each step, and achieves near the same accuracy.

6.3 Lawyers

The raw lawyer records follow much of the same deficiencies in quality as the assignee records. Again, we only offer a preliminary disambiguation of lawyer records using the same algorithm as described above, but future development will yield more accurate results.

6.4 Inventors

We provide a polished disambiguation mechanism for inventor records. Using the published name of an inventor, the patent technology class, co-inventor names, published location and original assignee, we are able to infer with more than 95% accuracy which inventor records are the same across all records in the patent database.

A detailed summary of our technique can be found through a related Fung Institute publication [11].

7 Statistics

Many research applications of patent data require records from multiple tables to be linked together: for instance, finding all citations made to a patent, or finding all patents for an inventor. Due to the size of the database, however, gathering all the requisite data and linking it together takes a nontrivial amount of time. To facilitate some common research vectors, we provide three tables of precompiled statistics.

The **FutureCitationRank** table contains the rank of each patent by the number of future citations in each year. This answers the question “in year X, patent number Y got Z citations. It was the Nth most cited patent that year”.

The **InventorRank** table contains the rank of each inventor by how many patents they have been granted in a given year.

The **CitedBy** table contains the direct mapping of a focal patent to all patents that cite that patent.

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