A KNOWLEDGEBASE OF STRESS REPONSIVE GENE REGULATORY ELEMENTS IN *Arabidopsis thaliana*

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Curation

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Transcription factors

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Stress response

ABSTRACT

Stress responsive genes play a key role in shaping the manner in which plants process and respond to environmental stress. Their gene products are linked to DNA transcription and its consequent translation into a response product. However, whilst these genes play a significant role in manufacturing responses to stressful stimuli, transcription factors coordinate access to these genes, specifically by accessing a gene's promoter region which houses transcription factor binding sites. Here transcriptional elements play a key role in mediating responses to environmental stress where each transcription factor binding site may constitute a potential response to a stress signal.

Arabidopsis thaliana, a model organism, can be used to identify the mechanism of how transcription factors shape a plant's survival in a stressful environment.

Whilst there are numerous plant stress research groups, globally there is a shortage of publicly available stress responsive gene databases. In addition a number of previous databases such as the Generation Challenge Programme's comparative plant stressresponsive gene catalogue, Stresslink and DRASTIC have become defunct whilst others have stagnated.

There is currently a single *Arabidopsis thaliana* stress response database called STIFDB which was launched in 2008 and only covers abiotic stresses as handled by major abiotic stress responsive transcription factor families. Its data was sourced from microarray expression databases, contains numerous omissions as well as numerous erroneous entries and has not been updated since its inception.

The Dragon Arabidopsis Stress Transcription Factor database (DASTF) was developed in response to the current lack of stress response gene resources. A total of 2333 entries

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were downloaded from SWISSPROT, manually curated and imported into DASTF. The entries represent 424 transcription factor families. Each entry has a corresponding SWISSPROT, ENTREZ GENBANK and TAIR accession number. The 5' untranslated regions (UTR) of 417 families were scanned against TRANSFAC's binding site catalogue to identify binding sites.

The relational database consists of two tables, namely a transcription factor table and a transcription factor family table called DASTF_TF and TF_Family respectively.

Using a two-tier client-server architecture, a webserver was built with PHP, APACHE and MYSQL and the data was loaded into these tables with a PYTHON script. The DASTF database contains 60 entries which correspond to biotic stress and 167 correspond to abiotic stress while 2106 respond to biotic and/or abiotic stress.

Users can search the database using text, family, chromosome and stress type search options. Online tools have been integrated into the DASTF database, such as HMMER, CLUSTALW, BLAST and HYDROCALCULATOR. User's can upload sequences to identify which transcription factor family their sequences belong to by using HMMER. The website can be accessed at http://apps.sanbi.ac.za/dastf/ and two updates per year are envisaged.

DECLARATION

DECLARATION

I declare that "A KNWOLEDGEBASE OF STRESS REPONSIVE GENE REGULATORY ELEMENTS IN *Arabidopsis thaliana* "is my own work, that it has not been submitted for degree or examination at any other university, and that all the sources I have used or quoted, and all work which was the result of joint effort, have been indicated and acknowledged by complete references.

MUHAMMED SALEEM ADAM MAY 2011

Signed: _______________________________________

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CHAPTER ONE

INTRODUCTION

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INTRODUCTION

A plant's response to the encroachment of a wide range of environmental stresses is mediated by its sessile nature which essentially shapes its ability to respond to these factors. These demands which negatively affect a plant's metabolic functioning, growth and adaptive capacity are referred to as stress. Stress may thus be viewed as a condition effected by a stressor (singularly or in combination) leading to a stress response which may result in damage such as cell death, organ incapacitation and permanent tissue and organ damage (Lichtenthaler 1995). Furthermore, a plant's response to stress-related events tacitly assumes a physiological norm. That is, a basal level implying environmental conditions supplying required quantities of water, light, temperature and nutrients under which a plant is able to thrive. An occurrence of stressors or stress events above a particular threshold or physiological norm would thus constitute a significant deviation and hence activates the need for a significant response. For example, a substantial increase in the intensity of environmental temperature requires a consequent change in the organism's perception and classification of the respective heat-related stimuli. That is, the level of stress signaling as indicated by responsive signal transduction mechanisms has exceeded a basal limit to a point where the increase in demand intensity can no longer be compensated for by conventional responses. At this point, stress is no longer perceived as something competitive but as hostile and is processed as an attack which requires appropriate defence operators and mechanisms (Lichtenthaler 1998). At this point a distinction should be made between biotic and abiotic stress. Stress events such as heat, cold, water deprivation, salinity, flooding, pesticides and soil nutrient depletion are classified as abiotic stress, that is, non-live stress

entities. Entities such as viruses, pathogens, necrophytes, biotrophs and insects are classified as biotic or live stressors that are capable of responding to a host's defence strategies.

1.1 TRANSCRIPTIONAL REGULATION IN PLANTS

Transcription is the process whereby response mechanisms coded in a native language or instruction set are operationalised in relation to environmental stress stimuli. The system operates via a network of reception, activation and amplification via signal transduction components where specific genes have been proposed to have stimuli-specific responses (Rodríguez, Canales and Borrás-Hidalgo 2005).

Many biological processes in a plant are regulated at the level of transcription, indicating the manner in which genes are expressed in relation to environmental stimuli (McGinley 2000). Changes in gene expression have been shown to underlie responses to environmental cues and stresses such as changes in light, temperature, nutrient availability and defence responses to pathogens (Aarts and Fiers 2003). The above responses are mirrored in an intricate network of components and mechanisms where signaling and transduction systems operate, activating/deactivating and shuttling response cues to various parts of the plant (Mahajan and Tuteja 2005). Furthermore, whilst the mechanisms of transcription are largely common across eukaryotes, their components vary among kingdoms (Arabidopsis Genome Initiative 2000; Riechmann et al., 2000).

Transcription factors are protein complexes that can shape the relationship between an organism and its response to environmental stress by influencing (initiate, enhance or inhibit) the transcription of specific genes. RNA polymerase is the enzyme that transcribes genes to make messenger RNA, which is used to generate the necessary response-proteins required by the system. Transcription factors assist RNA polymerase to bind to specific segments of DNA in an area known as the promoter region. The promoter is a regulatory region of DNA which marks the target site where RNA polymerase binds and also known as the transcription start site.

Every gene has a promoter region, however in the case of eukaryotes, its location may vary. For example, some promoters are located towards the three prime (3') region of the gene. The binding of RNA polymerase to the start site initiates the transcription process where instructions are written to the coding region of a gene*.* In eukaryotes, the promoters of many (but not all) genes contain the sequence TATAA, also known as the "TATA box". This region is twenty-five to thirty nucleotides upstream from the transcription start site. This sequence, in turn, is recognized by the TATA-binding protein (TBP) (Riechman 2002). The TBP binds to the sequence thereby marking the start site of transcription. By controlling RNA polymerase's access to the gene, transcription factors control the rate at which a gene is transcribed, thereby effectively regulating the rate at which genes are expressed. This control of the transcription process is underscored by the relationship between transcription factors and DNA cis-regulatory elements occurring upstream in the five prime untranslated region (5' UTR) of a gene. That is, a gene's responsiveness to certain stimuli is a result of their predisposition or 'hard wiring' to these cis-regulatory elements (Zhang et al., 2005).

However, experimental data on these binding specificities are scant and with respect to *Arabidopsis thaliana,* approximately three percent of these binding sites have been determined experimentally (Schröder et al., 2010). Furthermore, wet lab based

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determination of transcription factor binding sites (TFBS) using traditional DNA footprinting and chromatin immunoprecipitation (ChIP) technologies such as CHIPsequencing (ChIP-seq) can be time consuming and expensive (Grau et al., 2005; Chan et al., 2011). Hence numerous binding site prediction tools such as MATCH (Kel et al., 2003), VOMBAT (Grau et al., 2005) and PROMOTERSWEEP (Val et al., 2009) have been developed.

1.2 TRANSCRIPTION FACTORS

Transcription factors operate in an environment sensitive fashion, contingent on the tissue or cell-type that they occur in and the environmental stimuli that trigger their activation. They possess discrete DNA binding domains which are functionally specific and are drawn from a limited set of motifs (Latchman 1997).

These domains mediate sequence specific binding features by recognizing and matching an attendant, compatible series of nucleotide bases. It is the latter selective DNA recognition process and hence response generation process which underlies the cuespecific responses of an organism (Riechmann and Ratcliffe 2000). That is, they generate sequence specific control over transcription factor bindings thereby eliciting timeous, context sensitive and response specific results. It is in this context that the transcription factor binding sites are referred to as response elements. DNA binding domains (DBD) are used to either bind directly to DNA or as part of a large protein complex. DNA binding domains possess structural motifs or recurring elements which are used to divide transcription factors into classes (also known as superclass), families and subfamilies (Riechmann et al., 2000). Furthermore, individual family or sub-family members may play contrasting roles. For example, some members may play an activating role whilst others play repressive roles. Based on the structural motif concept, five main structural classes or superclasses can be identified namely, basic domain Helix-turn-Helix domain, zinc coordinating domain, beta scaffold with minor groove contacts domain and other domains (Table 1.1) (Stegmaier, Kel and Wingender 2004).

Transcription factor families which have been implicated in *Arabidopsis thaliana* stress response activities are AP2-EREBP, BZIP, bHLH, HSF, MYB and MYB-related, NAC and WRKY (Glazebrook 1999; Singh, Foley and Onate-Sánchez 2002; Eulgem 2005; Eulgem et al., 2005; Varshney and Koebner 2007; Bu et al., 2008; Van Verk, Gatz and Linthorst 2009).

Transcription factors also use signal transduction pathways as messaging mechanisms to respond to stress related signals, specifically, pathways related to the hormones salicylic acid (SA), jasmonic acid (JA) and ethylene (ET) (Numchuk et al., 2003; Kwon 2010). Ethylene is involved in various developmental processes, such as plant growth and fruit ripening. Besides these processes, ethylene is also involved in environmental stress signaling upon wounding or pathogen attack. Jasmonic acid is induced in defence related activities in response to necrotrophic pathogens and herbivorous insects whilst salicylic acid plays a central role in recognizing pathogen signatures and is induced after attack by biotrophic pathogens (Dong et al., 1991; Chung et al., 2008).

Furthermore, accompanying the increasing body of literature reporting on stress responses in plants, is a plethora of genes encoding biotic and abiotic stress-related genes (Hirt and Shinozaki 2004; Jenks and Hassegawa 2006; Rao, Raghavendra and Reddy

2006; Hirayama and Shinozaki 2010; Pareek et al., 2010). Hence the need for a database which collects and integrates this information.

1.2.1 TRANSCRIPTION FACTOR DATABASES

There are *A*. *thaliana-*specific transcription factor databases such as DATF (Database of Arabidopsis Transcription Factors) and Athamap (Steffens et al., 2005) as well as plant transcription factor databases which have an *A. thaliana* sub-category. These include the German PlnTFDB (Plant Transcription Factor Database) (Perrez-Rodrıguez et al., 2010) the Chinese PlnTFDB (Plant Transcription Factor Database) (Zhang et al., 2011) and RARTF (RIKEN Arabidopsis Transcription Factor database) (Iida et al., 2005). These databases do not focus on stress related issues but on transcription factors and their families. The Plant Stress Gene Database is a cross-species database that has an *A. thaliana* sub-section but only contains 33 entries and does not have any associated **WESTERN CAPE** analytical tools.

There is currently one *A. thaliana* -specific stress related database namely, Stress Responsive Transcription Factor Database (STIFDB). It contains 2629 entries which have been sourced from public microarray related databases. Genes which were significantly upregulated in response to abiotic stresses such as cold, salinity, light and water were selected as candidates for their database and organized in transcription factor families (Shameer et al., 2008). STIFDB's methodology for its binding site prediction algorithm was based on identifying ten families known to be involved in abiotic stress response which in turn led to the creation of a set of 22 Hidden Markov models (HMM). The upstream sequence including the 5' UTR of each gene was scanned using these

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models to predict a binding site for each transcription factor. The database features the sequence alignment tool BLAST and has not been updated since its inception in 2008.

1.3 DATABASE DESIGN MODELLING PROCESS

A database design requires a context model that represents the most salient aspects of the problem. In other words, the context model serves as the starting point for the iterative identification of objects and relationships constituting the basis of an information model. The process starts with understanding a client's problem context with a view to eliciting information which serves as input to the development of a specification. This, in turn, is interpreted using data model-based conceptual and notational techniques (Avison and Fitzgerald 1995).

Information elicitation is an iterative process which involves continuous interaction with users and their target context. Information may be captured and conveyed by various means. Human language is the most important conveyor and repository of meaning. However, meanings, derived from language-use can be vague and ambiguous. Context can play an important role, leading to the interpretation of words in varied and contradictory ways. Information, to a certain extent, is also dependant on the user and their understanding of their needs. That is, analysts will glean information filtered through the client's interpretative lenses. Furthermore, a client is not a monolithic entity, but rather refers to the various role players belonging to the single problem context. Hence, there is a variety of perspectives which may emerge (from a single context) affecting the quality of the information produced (Satzinger, Burd, and Jackson 2002). The sources of information may vary from verbal descriptions to various documentary pieces of information describing the current context. Evolving from the process is the encapsulation of the information in terms of a model, specifically, a context model. The model serves as a 'concrete' basis for further iterative refinements but most importantly a base around which central objects and their relationships are identified (Easterbrook 1993).

In the bioinformatics community, Ontologies such as Gene Ontology (GO) have specifically been developed to minimize the problems related to communication and reference. GO terms ultimately aim at developing a standardized vocabulary whereby a common set of terms and their respective meanings can be used across the vast spectrum of biological researchers and their respective communities (Helden et al., 2000; Harris et al., 2004).

The context model (Figure 1.1) attempts to capture some of the basic aspects of transcription initiation-regulation. The stimuli processing model indicates the rudimentary path of stimuli reception via signal transduction mechanisms to the transcription factor machinery which is implicated in chromatin remodelling. This process opens a path to the promoter region of DNA which houses transcription factor binding sites. Once transcription factors bind to these sites it initiates the transcription process. Furthermore, a second path is related to certain transcription factors which, irrespective of the chromatin packaging are able to recognize and bind to their respective target binding sites. Hence the core objects under investigation can be identified, namely, transcription factors, binding sites, promoter, DNA and the relational aspect of transcription factors binding to binding sites. These objects operate in a biological context and as such are located in a network of other biological objects which assist in developing a more complete picture of the target context, namely, the biological information model (Figure 1.2).

The development of context model also initiates the model conversion process, whereby an initial non-technical model is gradually transformed into a database design. The information model (Avison and Fitzgerald 1995) basically identifies and abstracts biological objects to the level of a sequence concept, namely, a sequence of nucleotides and amino acids. Furthermore, it assists in identifying additional objects, which, in this instance is the aggregate or family object formed on the basis of identifiable motifs/domains or shared sequence or structural signatures. The model also introduces non-biological objects such as the annotation object which is predicated on GO terms and journal publications. The basic idea behind the annotation concept is that all information in this model is ultimately referenced by publications.

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Figure 1.1: Context model for transcription factor initiation

The context model captures some basic aspects of transcription initiation regulation. Firstly, the rudimentary path of stimuli reception via signal transduction mechanisms to the transcription factor machinery which is implicated in chromatin remodelling. Secondly, chromatin remodelling opens a path to the promoter region of DNA which houses transcription factor binding sites. Transcription factors bind to these sites and initiate the transcription process.

Figure 1.2: Biological information model

This model describes the basic information requirements for each relationship and its participants. For each transcription factor, a series of property and object relationship dimensions are produced.

1.4 THE CONCEPTUAL MODEL

There is a difference between the relational model and the entity-relational model. The relational model was developed by Codd in 1969 whilst the entity-relational model developed by Chen in 1976 (Silberschatz, Korth and Sudarshan 2005). The entityrelational model is a way of capturing/modeling a reality in terms of objects and their dependencies. It is amenable to a variety of contexts and as such has found particular use in systems analysis and design methodologies. This entity relational model, which uses entity relational diagrams (ERD) as a means of depiction, can then be led along different logical modeling paths including a path leading to Codd's relational model. That is, the ERD approach is used to diagrammatically map a logical path or argument to particular design structure at a particular level. In other words, it may be viewed as depicting the logical structure of the premises leading to a particular relational database design (Whitten and Bentley 2002). **UNIVERSITY** of the

The 'implementation' detail of the structure is then administered by applying the principles of the relational model. In terms of Codd's model, reality is described in terms of records expressed as a tabular structure consisting of rows and fields where fields are object properties or object relationship properties and rows are populated with instances of object values.

A conceptual model (Figure 1.3) attempts to map the entire background to the network of biologically related objects from the association between transcription factor-genebinding site relationships to the gene's production of a response protein. Here, the conceptual model has been appropriated to represent the biological flow of information whilst located within the background of molecular biology's 'Central Dogma' (Keet

2003). The latter constitutes the business rules which determine the 'objects of interest' and the rules which bind these 'objects of interest' as relationships (Nelson, Reisinger, and Henry 2003).

ERDs use a technique where entities and their properties represent the 'objects of interest' about which information is stored and relations depict the associative links between entities (Chen and Carlis 2003). The numerical additions indicate cardinality or the level of the relationship between entities. For example, entities which start with a number greater than zero indicate that they are conceptually, mandatory participants in a relationship. Furthermore, a number greater than one indicates the 'many' side aspect of the participant. A many to many relationship is depicted as 'M:N'. Essentially, it means that the relationship between participating entities have numerous associative permutations or are conceptually non-limited in terms of their combinations. For example, any gene can theoretically, produce numerous proteins and proteins in turn may be associated with numerous participating genes, thus leading to a many to many relationship (Bornberg-Bauer and Paton 2002).

Figure 1.3: A conceptual model of biological entities

A global view of entities and their relationships. Namely the relationship between entities such as species, organism, genome, chromosome, gene, promoter, transcription factor binding sites, transcription factors, proteins, motifs, domains and orthologs.

1.5 NORMALISATION

As the modeling process moves to the logical design stage, decomposition takes place. That is, objects and relationships are decomposed to discrete self-subsisting units. Specifically, the 'many to many' relationships are all dissolved into 'one to many' relationships. Strategically, the process may be viewed as the unbundling or dissection of the original target context into discrete units which can then be reconstituted in numerous ways. This process allows the user to generate information of the target context in novel and insightful ways which is a function of the database querying process.

There is a fundamental set of guidelines to this information structuring process which is referred to as the process of normalization (Atzeni et al., 1999). It is implemented via the usage of the so-called normal forms and guides the development of a consistent database which reduces redundancy and removes anomalous behaviour. Normal forms guide the building of objects in terms of distinctive properties which uniquely identifies a row of data. This is referred to as a primary key and is used to locate data and bind relationships which are now viewed as dependencies connecting objects or tables of data. During the querying process these keys are used by the database to join tables, make calculations and locate data. The normal forms are structured hierarchically, that is, graduation to a subsequent normal form requires compliance with the previous normal form. Hence, the second normal form requires compliance with the first normal form.

The first normal form requires management of the most basic aspect of database design. As indicated earlier, this involves the decomposition of objects into discrete atomic units such that a column or field represents a single property or unit of information. This also requires the decomposition of multi-valued attributes into single-valued attributes (Kent 1983).

The next and most important principle of the second normal requires that all properties referred to as non-key properties have to be dependent on the primary key. The primary key as indicated earlier essentially defines a row of data with a unique property or set of properties. That is, more than one column participates in key formation and as such it is called a composite key. Hence, non-complying non-key properties may point to information about a separate object which can be modeled using a separate table and in turn can then be constructed with its own primary key. In addition to compliance with the second normal form, the third normal form requires the mutual exclusivity of table non-key properties or columns. That is, whilst the second normal form requires the dependence of all non-key properties on the primary key, the third normal form requires that each non-key property should be mutually independent. Hence, any transitive dependencies between non-key properties should be identified and resolved

1.6 SWISSPROT

SWISSPROT is a curated protein database that aims to provide a high level of annotation specifically functional annotation (Figure 1.4). This data layout provides a useful starting point to retrieve quality data for plant stress-related genes. The SWISSPROT record contains amongst other fields, the "KEYWORD", "GO" and "COMMENTS" fields. Each of these fields can be used to carry out a comprehensive search of SWISSPROT. Keywords are used to provide a summary of a record's contents which is then indexed according to a set of ten categories. These include biological process, cellular component,

molecular function, coding sequence diversity, developmental stage, disease, domain, ligand and post-translation modification. GO terms and SWISSPROT keywords are stored under the same section namely, the "Ontologies" section of an entry (Figure 1.4) (Schneider et al., 2009). GO annotations are applied manually and keywords reflecting the contents of these GO annotations are also generated manually. GO annotations are classified in terms of evidence codes which are used to describe the source and strength of a particular annotation. For example, 'Inferred From Direct Assay' (IDA) indicates that the source for an annotation is an experiment and hence has the highest level of evidence classification. Whilst 'Inferred From Electronic Annotation' (IEA) is applied where an annotation has taken place though an automated computational procedure and a curator has not personally verified the annotation such as through automated importing of annotations from a related database. (Berardini et al., 2007).

Finally, the 'comments' section, which follows a similar logic to the 'keyword section', generally conveys information about a protein's function. Annotations in this area are grouped according to 'topics' such as Function, Induction, Involvement, Enzyme regulation, Pathway, Subcellular location, Tissue specificity and Developmental stage. Furthermore, as in the case with keywords, comments about a protein's function require the use of standardized terms to facilitate text searches and database interoperability (Schneider, Tognolli and Bairoch 2004; Scheider et al., 2009).

Due to its usage of manual curation SWISSPROT is regarded as a database with high quality data however its annotation cannot be accepted at face value. Even at the most basic level there is always the possibility of error. For example, the following entries**:** http://www.uniprot.org/uniprot/Q9M9V8

http://www.uniprot.org/uniprot/Q39016

contain the annotation comment "By drought and high-slat stress". The phrase 'high-slat

stress' should read 'high salt stress'**.**

General annotation (Comments)

FIGURE 1.4: The 'General annotation' section of a SWISSPROT record

The major categories are the 'Comments' and 'Ontologies' section. With regard to this entry, the GO 'Biological process' and 'Molecular function' category shows transcription factor involvement in cold acclimation via transcriptional regulation. The 'Function' category indicates the details of the stress response. The cited references confirm the annotations as correct. Hence the 'Keywords' section shows the terms 'stress response', 'transcription' and 'transcriptional regulation'.

1.7 MOTIVATION AND RATIONALE

Existing food shortages in tandem with changing weather patterns have exacerbated global food security especially in relation to staple food crops such as rice and maize.

In part, changing weather patterns provide the basis for environmental stresses as experienced by plants. An understanding of the mechanisms by which plants perceive and process responses to extremities, may provide insights which can be applied to developing stress resilient crops (Hirt and Shinozaki 2003).

The study of stress responsive genes is facilitated through using *Arabidopsis thaliana* as a point of reference due to its portability, availability, easy cultivation and comparatively small genome size. Furthermore, the availability and accessibility of information about *Arabidopsis thaliana* has also played a role in reinforcing its status as a model plant organism (Bevan and Walsh 2005).

Globally, the availability of numerous plant stress related research groups is not mirrored by a similar availability of public database resources for plant stress genes. This scarcity has been reported by the journal Nucleic Acid Research (NAR) (http://www.oxfordjournals.org/nar/database/subcat/13/39). The few databases that are still available such as the Plant environmental stress transcript database, the Plant Stress Gene database and the Stress Genomics database have not been updated since 2006 whilst others such as the Generation Challenge Programme comparative plant stressresponsive gene catalogue (GCP) have long since become defunct (Balaji et al., 2006; Wanchana et al., 2008). Furthermore, the Plant Stress Gene database and the Stress Genomics database do not have any accompanying academic publications and hence it is difficult to assess their quality.

The existing database of abiotic stress related transcription factors, STIFDB, unfortunately suffers from a number of problems. Firstly, in terms of the exclusion of stress related genes, some basic abiotic stress related transcription factors have been omitted from STIFDB (Table 1.2). Secondly, transcription factors have been included that have no correlation to abiotic stress (Table 1.3). For example, three of the entries have no abiotic stress related function. In fact, their functional annotation as indicated by the 'Stress Response' column is incomplete and the PATHOGENESIS-RELATED GENE 1 is involved in biotic stress-related defence.

Due to the key role transcription factors play in plant responses to environmental stress, the current project Dragon Arabidopsis Stress Transcription Factor database (DASTF) hopes to fill a gap in stress related transcription factor databases. The development of a manually curated database will assist in structuring the understanding of these stress responsive mechanisms. In turn, reliable data may serve as a springboard for harnessing analytical and predictive tools to generate greater insight into the environmental response mechanisms of *Arabidopsis thaliana*.

Domain superclass	TF family
Basic domain	BES1, bHLH, bZIP, EIL, GeBP, TCP
Helix-turn-helix domain	ARR-B, E2F-DP, FHA, G2-like, HB,
	HSF, MYB, MYB-related, WP-RK, Sigma70-
	like, zf-HD
Zinc coordinating domain	Alfin-like, C2C2-CO-like, C2C2-Dof, C2C2-
	GATA.
	C2C2-YABBY, C2H2, C3H, CPP, GRF,
	HRT, LIM,
	PHD, PLATZ, SBP, SRS, TAZ, VOZ,
	WRKY, ZIM
Beta-scaffold with minor	CCAAT, CSD, GRAS, HMG, MADS
groove contacts domain	
Others	AP2-EREBP, ARF, ARID, BBR/BPC,
	CAPE CAMTA, DBP,
	DDT, Jumonji, LFY, NAC, NOZZLE, PBF-
	2-like, RB,
	S1Fa-like, Trihelix, TUB, ULT, ABI3VP1

TABLE 1.1: Plant transcription factor families and their DNA binding domains (Panchon 2008)

TABLE 1.2: Abiotic stress-related transcription factors omitted from STIFDB

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TABLE 1.3: Erroneous entries included in STIFDB

1.8 AIMS AND OBJECTIVES

- 1. Produce a manually curated set of plant stress-related proteins
- 2 .Identify transcription factor binding sites for the above proteins
- 3. Develop a web portal for retrieving plant stress-related regulatory information

1.9 THESIS OUTLINE

This thesis consists of four chapters:

Chapter 1:

• Brief overview of (i) stress-related genes in plants, (ii) transcription regulation,

(iii) current databases for stress-related genes in plants and (iv) database design

Chapter 2:

 • Describes the methodology for collection and curating stress-related genes and webserver development. ESTERN CAPE

Chapter 3:

• Describes the data captured in the DASTF database and the functional

features.

strategies

Chapter 4:

• Discussion and Conclusion

CHAPTER 2

METHODS

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2.1 COLLECTION OF STRESS RESPONSE TRANSCRIPTION FACTORS

Using the SWISSPROT query engine, three text-based searches were carried out to retrieve regulatory elements that respond to environmental stress as follows**:**

- **(a)** organism:"*Arabidopsis thaliana* [3702]" AND keyword:"Stress response [KW-0346]"
- **(b)** organism:"*Arabidopsis thaliana* [3702]" AND go:"response to stress [0006950]"
- **(c)** organism:"*Arabidopsis thaliana* [3702]" AND annotation:(type:function stress)

The above queries generated 209, 2849 and 353 records respectively. Duplicated records were removed and resulted in final dataset of 2904 entries specific for stress response proteins.

A total of 2333 out of 2904 SWISSPROT records were mapped to their corresponding GENBANK and The Arabidopsis Information Resource (TAIR) IDs using a SWISSPROT online tool called "ID-Mapping Tool" (http://www.uniprot.org). A PYTHON parser script was written to load the 2333 records into a local MYSQL database (see section 2.4 for database design).

2.2 CURATION

The adopted curation strategy was based on experience with a project called DAMPD (Sundarajan et al., in prep.) which is a collection of manually curated antimicrobial peptides. The curation methodology involved checking SWISSPROT keyword terms and function annotation (in the 'general annotation (comments)') against the references listed for each record. That is, if the SWISSPROT keyword section had the terms "response to stress", it was checked against the function annotation section and then cross-referenced
against the list of sources for the respective entry. Each record was curated manually using this procedure. SWISSPROT protein existence codes or evidence at protein level (PE codes) were also appropriated in the same manner namely PE level one (evidence at protein level) and PE level two. PE level one indicates that there is clear experimental evidence for the protein's existence. PE level two indicates that although there is no clear experimental evidence, there is at least gene expression data pointing to the existence of a transcript. Finally, two additional stress response databases namely STIFDB and Plant Stress Gene Database were cross-referenced to retrieve further information where applicable.

2.3 WEBSERVER DESIGN

The system was implemented using a tiered client-server architecture where an application is essentially divided into three layers, namely, a presentation layer, a logic layer and a data layer. The organized grouping of these layers as in the case of a two-tier or three-tier system determines the type of client-server architecture (Ramanathan 1995). The DASTF system was implemented using a two-tier architecture which is also known as a thin client because the majority of processing is handled by the server (Figure 2.1). These include the logic and data layers whilst the client only handles the presentation and layout processing (Steiert 1998).

The presentation layer deals with the user interface of the client machine and it provides the end-user with the visual means of accessing and querying the system. This layer handles the resulting output of a request which is formatted in Hypertext Markup Language (HTML) using cascading style sheets as a means to manage the appearance of a webpage.

The logic layer constitutes the so-called business rules of the application and is responsible for error and conformance checking. That is, making sure the user has filled in the necessary details as provided by the web interface which in turn constitutes the parameters of a request (Ward and Dafoulas 2006). As part of the logic layer, the latter process continues by translating the client request using its business logic modules into a database readable format, namely, Structured Query Language (SQL). The database was designed using MYSQL (Figure 2.1) and resides on an Apache webserver that acts as a medium routing requests from the client to the database. In other words, PHP uses its inbuilt modules to speak to the database via the Apache software.

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Finally, the data layer is made up of the database connection layer and the database layer. The data connection layer is responsible for connecting a user request to the actual database (Ward and Dafoulas 2006). It functions as a generic component capable of connecting to a database irrespective of the source database's architecture (network, hierarchical or relational), location path (where the database resides) or vendor (Oracle, Microsoft, Borland, MYSQL). In turn, the database layer processes (inserts, updates, deletes, queries) the request in terms of its own set of constraints, such as integrity checking for example. It then returns the result of the request along the same pathway it was received (Bahrami 1999).

2.4 DATABASE DESIGN

Two MYSQL tables were generated namely DASTF_TF and TF_Family tables (Figure 2.2). Each SWISSPROT record was parsed using a PYTHON script to extract fields needed to populate the MYSQL database (Appendix I).

2.5 STRESS RESPONSE TRANSCRIPTION FACTOR FAMILIES

Stress response proteins were mapped to ENSEMBL IDs using a SWISSPROT IDmapping tool and each ENSEMBL ID was used to extract 5'UTRs from the ENSEMBL *Arabidopsis thaliana* core database. The 5' UTRs were scanned for transcription factor binding sites with TRANSFAC professional database version 2011.1 and the output file was parsed with a PERL script (Appendix II) (Matys et al., 2006).

Stress response genes were divided into two exclusive type groups, namely those which only respond to abiotic stress and those which only respond to biotic stress.

The terms plant defence, bacteria, fungus, virus, pathogen, nematode, oomycete, wounding, insect, chitin, jasmonic acid, ethylene and salicylic acid were used to identify biotic stress. Abiotic stress was identified by terms related to temperature (cold, heat), light (ultraviolet, red light, blue light), water (drought, flooding, deprivation), precipitations (hail, snow, frost, fire), abscisic acid, salinity, herbicide and pesticide. The exclusive categories were tested for any combination of these terms. Those which had any combination of these terms were excluded from that category and placed into a 'biotic and or abiotic' category.

Figure 2.1: The two-tier client-server model implemented in DASTF

The majority of processing takes place on the second tier. In this implementation the second tier refers to the server side also known as 'back-end' of the system. The client has minimal processing load and is also known as the 'front-end' of the system.

Figure 2.2: Entities with their associated properties

 Transcription factor details are stored in the DAST_TF table and includes it's name, accession number, sequence and associated GO descriptions. Transcription factor family data is stored in the TF_Family table. Such as the family name, transcription factor members of the family and their binding sites.

CHAPTER 3

RESULTS

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3.1 DASTF WEBSERVER

The DASTF system was developed with a two-tier client-server architecture which is better suited to smaller projects where a faster application development time is required and in an environment where user traffic is expected to be lower (Hemmer 1995). Hence it is suited to a small community-specific project such as DASTF where batch user downloads are not supported.

The DASTF database (Figure 3.1) currently holds 2333 entries. SWISSPROT's 'ID mapping' tool was used to trace these entries to their TAIR and ENTREZ GENBANK counterparts. These genes cover both biotic and abiotic stresses such as cold, heat, light, salt, water deprivation, wounding, oxidation, fungi and bacteria. There are two query categories, namely, an exclusive category and a related category. Genes that are found in the exclusive categories are either exclusively biotic or exclusively abiotic which means that there is no overlap in stress response between these genes. A total of 60 entries only correspond to biotic stress whilst 167 entries only correspond to abiotic stress. The related type stress category refers to genes that respond to biotic and/or abiotic stress types. The overwhelming majority are found in this category namely 2106 entries.

Furthermore, there are 424 transcription factor families of which 417 contain genes for which UTRs (untranslated region) could be extracted from ENSEMBL and UTRs shorter than 50 nucleotides were excluded.

Unique transcription factor binding sites were identified in 13 stress response genes belonging to characterised protein families (Table 1.4). A total of 10 stress response genes without any protein family classification contained unique transcription factor binding sites. Among these unique binding sites was SED motif that bound to both

AT2G42910 (methyltransferase superfamily) and AT3G12810 (unclassified protein family).

Chromosome one, five, three, two and four contain 599, 574, 417, 372 and 371 genes respectively. The most common abiotic related stresses in the database are responses to light, heat, salinity, cold, water deprivation and oxidative stress respectively. The distribution of abiotic only and biotic only responsive genes mirrors the overall distribution of stress responsive genes over the five chromosomes.

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Table 1.4: Unique TFBS motifs in stress-related genes families

3.3 WEBSITE OVERVIEW

The DASTF database can be accessed via the 'search page' (Figure 3.2). At present, there are four search options, namely, a simple search, search by stress type, search by chromosome and search by transcription factor family. The stress type search allows the user to search by two categories, namely, an exclusive stress-type response and a related type stress response. The exclusive option consists of two types, that is, 'BIOTIC' or 'ABIOTIC' which is followed by the 'BIOTIC AND OR ABIOTIC' category.

The text search can be used in instances where the user has a protein name or a list of accession numbers from UNIPROT, TAIR, ENTREZ or PFAM. The chromosome search option reports all stress-related genes on a particular chromosome. Transcription factor family search allows the user to search a set of transcription factor families encoding stress-related functions.

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3.4 DATA ENTRY RECORD

 Each entry is identified by a series of fields (Figure 3.3). The DASTF entry field is the database's unique accession number for each record whilst 'Entry name' identifies the name of the transcription factor. The 'Uniprot Accession', 'Gene ID' and 'TAIR ID' fields are the respective UNIPROT protein, ENTREZ gene and TAIR accession numbers. Links to UNIPROT, ENTREZ gene and TAIR accession numbers allow the user to crossreference annotations as well as to access more detailed descriptions from the respective source databases. Similarly, the 'Pubmed field' provides a link to the publications which reference a particular entry. The 'Family field' identifies the transcription factor family that maps to a protein. The 'KEGG' field allows the user to access pathways that intersect a transcription factor while the 'PFAM' field provides protein domain structures for a specific transcription factor. The 'PE' field identifies the evidence level used during curation to validate the protein. 'Evidence at protein level' or 'PE $= 1$ ' indicates that a protein was identified by experimental evidence. The Chromosome field describes the chromosomal location of a gene in base pairs. Furthermore, by clicking the chromosome hyperlink an image is generated showing the exact location of the gene as well as its proximity to other genes on the same chromosome. The 'TFBS' field is linked to a gene's transcription factor binding site and when a user clicks the 'Click here to see the TFBS' hyperlink a binding site (Figure 3.4) is reported for the particular gene.

The GO field provides the functional annotation for each record. Each annotation such as stress response, evidence and other related activities are referenced by their respective GO identifier. Finally, the sequence field shows the protein sequence.

3.4 TOOLS

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The database features four tools which are accessible via the tools page (Figure 3.4). These are BLAST (Altschul et al., 1990), CLUSTALW (Thompson, Higgins and Gibson 1994), HMMER (Eddy et al., 1995) and HYDROCALCULATOR (Tossi et al., 2002). For BLAST the user uploads a sequence or list of sequences to test against the database. The sequences have to be in a specific format called the FASTA format. The utility then checks the sequences against the DASTF database and generates a score indicating which sequences are most similar to the user's input sequences. This allows users to identify whether their sequences are potential stress related genes. Another approach is using HMMER where a model is generated by inputting sequences with a high degree of validity. That is, by using sequences where the entries have a high GO evidence type and a high SWISSPROT protein existence level. Protein sequences are subsequently searched against the model to assign them to families by using similarity as a basis for comparison. In this manner users can predict which stress related transcription factor family their sequences belong to. HYDROCALCULATOR generates protein analysis by analyzing amino acid hydrophobicity using scales such as the combined consensus scale (CSS), Kyte and Doolittle and Eisenberg.

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FIGURE 3.1: The DASTF Homepage

The homepage of the DASTF website. The menu section at the top allows access to the search engine and online tools.

Search options are divided into 'simple search', search by 'chromosome', 'family and search using 'stress' categories. The simple search uses entry names and accession numbers as input and the 'stress search' uses 'BIOTIC', 'ABIOTIC' and 'BIOTIC AND OR ABIOTIC' categories.

FIGURE 3.3: List of properties for entry DASTF_2241

The TFBS detail (circled in red) is hyperlinked and explained in Figure 3.4.

FIGURE 3.4: TFBS prediction associated with UTR for entry DASTF_2241

 'motif name' refers to the TFBS name in TRANSFAC; 'position' refers to the base position in the UTR; 'strand' indicates the positive (+) or negative (-) DNA strand; 'core score' and 'matrix score' are thresholds for binding efficiencies when using TRANSFAC; 'binding site sequence' is the motif identified by TRANSFAC.

FIGURE 3.5: Tools

A list of the online tools available BLAST, CLUSTALW, HMMER and HYDROCALCULATOR

CHAPTER FOUR

DISCUSSION

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4.1 METHODOLOGICAL ISSUES

The DASTF database contains 2333 *Arabidopsis thaliana* stress responsive genes that were collected from SWISSPROT, they were curated, organised into protein families and TFBS predictions were generated for the corresponding UTRs. A competing database, STIFDB published in 2008, relied exclusively on microarray databases to identify genes which respond to abiotic stresses such as cold, drought, salinity and light. Candidate genes had to be significantly upregulated in at least three replicate microarray experiments. Limitations of microarray experiments include repeatability of results, the influence of statistical methods when interpreting the data and the influence of environmental and background noise on results (Draghici et al., 2006). The GO consortium has advocated a set of cautionary guidelines where expression data is used to assign functions to genes and the evidence code 'Inferred from Expression Pattern' has $UNIVERSITY$ of the been developed for this purpose (Evidence Codes Group 2007). Firstly, their guidelines specifically state that it may be difficult to conclusively identify the function of a gene based on the timing of its expression pattern in relation to an experimental condition such as a stress.

Secondly, microarray expression data should not be used to assign GO 'molecular function' annotation claims. For example, genes that are upregulated during a stress response should rather be classified according to a GO biological process category such as 'response to stress' and not according to the 'molecular function' category (Evidence Codes Group 2007). The wisdom of these guidelines became apparent when STIFDB's microarray sourced data was reviewed. A number of the entries were reviewed manually by comparing them with annotations in source databases such as SWISSPROT, TAIR and GENBANK to identify any links with stress response. It was found that a number of genes (Table 1.5, Appendix III) with unknown functions were incorrectly included in STIFDB database.

The records for these genes with unknown function annotations were updated in 2010 and 2011 by GENBANK, SWISSPROT and TAIR. STIFDB was launched in 2008 which implies that inspite of a two year gap since the publication of its data, none of the updates to these gene records in GENBANK, SWISSPROT and TAIR indicate any relationship with stress response. Furthermore there is no indication that the STIFDB team had curated its data prior to its storage (Shameer et al., 2008).

The approach to gathering data for the DASTF database was influenced by the results of the STIFDB data review process and hence the need for a foundation based on well annotated and curated entries was identified as a priority.

The DASTF database covers both biotic and abiotic stresses whilst STIFDB only contains entries related to abiotic stress. Furthermore, STIFDB restricted their database to a list of 22 abiotic stress related families. However, abiotic stress responses are not limited to these 22 families and hence DASTF includes other abiotic stress related families such as glutathione peroxidase family, carotenoid oxygenase family, cytochrome P450 family, GST superfamily and a number of unclassified families as well.

Similarly, STIFDB predicted transcription factor binding sites for genes belonging to its set of 22 families whilst DASTF contains binding sites for biotic, abiotic as well as unknown families of stress response genes. Unique transcription factor binding sites were identified in 13 stress response genes belonging to characterised protein families and 10 stress response genes without any protein family classification. These motifs have been

implicated in stress regulatory environments involving responses to salt, oxidation, cold, abscisic acid and light (Singh 1998; Schwechheimer et al., 1998; Choi et al., 2000; Hasegawa et al., 2000; Jackoby et al., 2000; Grover et al., 2001; Memelink et al., 2001; Pastori and Foyer 2002; Shen, Cao and Wang 2008). Furthermore, for each gene that is identified, the database provides information identifying both the metabolic processes that a gene is involved in and as well as its location relative to other proximate genes. In addition, the database is further enhanced by the integration of tools such as HMMER, BLAST, HYDROCALCULATOR and CLUSTALW. An area which requires attention is the categorization of previously unclassified protein sequences into families. By using the HMMER tool proteins such as DNA repair protein REV1 and F16G20.140 have been identified as potentially related to the family '6-phosphogluconate dehydrogenase'**.**

4.2 COMPLEXITIES, LIMITATIONS AND FUTURE WORK

Arabidopsis thaliana has a smaller more compact genome in comparison with other organisms such as rice (Karlowski et al., 2003). The distribution of genes across *A.thaliana* five chromosomes reflects the need for compact organisation and economical usage of its genetic resources (Mayer et al., 1999; Arabidopsis Genome Initiative 2000; Holtorf, Guitton and Reski 2002).

It was reported (in chapter three) that 2106 genes out of 2333 stress responsive genes, were able to respond to both biotic and abiotic stress which may be interpreted in terms of a phenomenon called cross-talk. Genes use common stress related signalling pathways to generate complex cascades of gene expression in response to environmental stress.

For example, plant defence genes use signal transduction pathways such as jasmonic acid, ethylene and salicyclic acid to cope with a variety of viral, bacterial and fungal pathogens. Various permutations of pathway and transcription factor combinations impact on the activation and deactivation of specific types of biotic defence related genes (Fujital et. al., 2006; Century Reuber and Ratcliffe 2008). Cross-talk permutations have been postulated as a means whereby genes are able to fine-tune their responses to a wider range of threats from its environment. Furthermore, pathways are also able to operate independent from each other, collaboratively as well as antagonistically (Nimchuk et al., 2003; Van Verk and Gatz 2009).

The issue of context adds another layer of complexity to this study which influences the manner in which developmental processes can be viewed. In normal circumstances, various parts of an organism are supplied with nutrients on a systematic basis leading to its growth and development. However, in the event of a biotic attack as in the case of a virus, for example, a plant may voluntarily cease supplying the affected area with nutrients. That is, in order to contain and prevent the spread of the virus which needs nutrients to survive, the plant may effectively 'kill off' the affected part and starve the virus of nutrients (Lichtenthaler, 1995; Lichtenthaler, 1998). Hence annotations which identify the manner in which developmental processes such as nutrient transport and other context specific behaviour become important. From the perspective of database annotation, identifying context, the components involved and their context specific expression is one of the major limitations of the database. However, it should be noted that whilst the latter processes are inherently complex, its absence does not detract from the value of DASTF as a resource which provides a comprehensive, curated catalogue of stress response genes in *Arabidopsis thaliana***.**

Furthermore, the database needs to be evaluated and updated on a regular basis by identifying new annotations which can embellish and augment existing data thereby providing a more comprehensive view of stress related genes and proteins.

Areas that require attention are:

i) the construction of a systematic and comprehensive library of motifs and their stress responsive associations which can be used to classify new entries.

ii) there is an urgent need to classify the large number of currently unclassified transcription factor families in the database.

iii) additional levels of detail should be added to entries as in the case of stress responses to viruses, bacteria and fungi such as data detailing which types of fungal, bacterial and viral strains have been studied should be added to the database to provide a greater degree of specificity to its annotation.

Finally, the database can be extended to include other species such as rice and maize and thereby also attempt to identify orthologs between these species.

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APPENDICES

APPENDIX I

Python SWISSPROT parser script

Parses a SWISSPROT flat file into individual records and imports the records into

a MYSQL table called DASTF

```
#!/usr/lib/python2.4 
from  future import division # it ensures that the division 1/2 --> 0.5 and not 0
""
```
FILE: swiss.py USAGE: swiss.py flatfile ""

"" "

This script is for getting certains fields from UniProt flatfiles of a given accession """

<u>Mahahahahaha</u>

import random, math, sys, os, glob, time, numpy, MySQLdb

```
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def parse(swiss_file): 
          filename = open(swiss_file) \n<b>TERN</b> <i>CAPE</i>counter_RX = 0counter CC = 0label = []\text{seq} = \text{""}uni_list = [] for line in filename.readlines(): 
           field = line.split()if field[0] == "ID":
               print "ID = ", field[1]
                uni_list.append(field[1]) 
               ID = field[1]elif field[0] == "AC":
               print "AC =", field[1].split(";")[0]
                uni_list.append(field[1].split(";")[0]) 
           elif field[0] == "RX":
               counter_RX = counter_RX + 1if counter RX == 1:
```

```
 for i in range(len(field)): 
         if "PubMed" in field[i]: 
         pb = field[i].split("=")[1].split(";")[0] uni_list.append(pb) 
       print "Pubmed = ", pb 
       #print uni_list 
 elif field[0] == "DR":
      if field[1].split(";")[0] == "GeneID":
         gene = field[2].replace(";"," "") uni_list.append(gene) 
         print "GeneID = ", gene
         #print uni_list 
 elif field[0] == "PE":
       pe = line.split(":")[-1].replace(";", "").replace("Evidence", 
             "Evidence").rstrip() 
          print "PE = ", pe
          uni list.append(pe)
       #print uni_list 
 elif field[0] == "CC" and len(field) > 2:
     if field[2].split(":")[0] == "FUNCTION":
      field[3:] = [''.join(field[3:])] print "Function = ", field[3].split(".")[0] 
       uni_list.append(field[3].split(".")[0]) 
       #print uni_list 
     elif field[2].split(":")[0] == "SIMILARITY":
      counter\_CC = counter\_CC + 1if counter_CC = \frac{W}{1}: STERN CAPE
        #print field[3:] = [''.join-field[3:]]field[3:] = [''.join(field[3:])]print "Family = ", field[3].split(".")[0]
         uni_list.append(field[3].split(".")[0]) 
         #print uni_list 
  #if field[0] not in label: 
  # label.append[field[0]] 
  #if field[0] not in label: 
  # label.append(field[0]) 
  if field[0] not in ['ID', 'AC', 'DT', 'DE', 'GN', 'OS', 'OC', 'OX', 'RN', 'RP', 
      'RC', 'RX', 'RA', 'RT', 'RL', 'CC', 'DR', 'PE', 'KW', 'FT', 'SQ','//']: 
     field[0:] = [''.join(field[0:]))seq = seq + field[0].replace(" ", "") #print seq 
 print "sequence = ", seq 
 uni_list.append(seq) 
conn = MySQLdbconnet (db = "dastf")cursor = conn.cursor ()
```

```
 cursor.execute('insert into uniprot values
```
 ("%s","%s","%s","%s","%s","%s","%s","%s")'% (uni_list[0],uni_list[1],uni_list[2],uni_list[3],uni_list[4], $uni_list[5]$, $uni_list[6]$, $uni_list[7]$, $()$

def main():

```
if len(sys.argv) != 2:
         print 'Usage: python swiss.py swiss_flat_file' 
         sys.exit() 
swiss_file = sys.argv[1]
 parse(swiss_file)
```
 $if _name__ == ' _main__$: main()

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APPENDIX II

Perl TRANSFAC parser script

Parses TRANSFAC file containing binding site motifs and retrieves: TAIRID, Motif

name, Motif sequence, Strand direction, Position, Core score and Matrix score

```
#! /usr/bin/perl 
# 
# 
use strict; 
$/="\nScanning sequence ID:"; 
my $f = shift;my \text{Spec} = 0;
my %hash; 
open(F, $f); 
while (<b>F</b>) {
        $rec++;THE THE THE
         chomp; 
         next unless (\text{Spec} > 1);
          next if ($_=~/No sites found for this sequence/); 
        \{\$ =~/^(\s+)(\S+)/;
        my sacc = $2;
                                       UNIVERSITY of the
        # \, \text{Shash} \{ \text{Sacc} \} = 1;WESTERN CAPE
# print "$acc\n"; 
        my \omega lines = split(\Lambdan/);
          foreach my $line(@lines) { 
                 next if (\text{Blue} = \text{--} / \text{--} \text{--} \text{--} \text{--} \text{--} \text{--}next unless (\text{Rine} = \frac{1}{2} /^P/);
                   #print "$line\n"; 
                   my ($tfbs_id, $pos, $str, $cmatch, $mmatch, $string, $tfbs_name) = 
split(<math>\Lambda</math>s+/, <math>fline</math>);#print "$acc $tfbs_id $pos $str $cmatch $mmatch $string $tfbs_name\n";
                  my \omegae = ($pos,$cmatch,$mmatch,$str, $tfbs_name,$string);
                   push @{$hash{$acc}}, [@e]; 
          } 
} 
close(F); 
foreach my $acc(keys %hash) { 
        foreach my \mathcal{S}e(\mathcal{Q} \{\mathsf{Shash}\{\mathsf{Sacc}\}\}) print "$acc\t".$e->[0]."\t".$e->[1]."\t".$e->[2]."\t".$e->[3]." ".$e-
>[4]."\t".$e->[5]."\n"; 
          } 
}
```
APPENDIX III

TABLE 1.5: Extended list of erroneous entries included in STIFDB

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