

TUTORIAL 1: About GDR/ Getting Started

1. The GDR homepage can be reached by going to <http://www.rosaceae.org>

GDR | Genome Database for Rosaceae

General Info | Species | Projects | Maps | Search | Tools | Community | Forums | myGDR

site search go
Home | Contact | About

GDR Quick Start

- Disclaimer
- About GDR
- Feedback Form
- Data Overview
- GDR Tutorial
- GDR Newsletters
- GDR Outreach
- Rosaceae Community

GDR Species

- Apple Pear
- Prunus Almond
- Apricot Cherry
- Peach Raspberry
- Rose Strawberry

Maps and Markers

- GDR Maps
- Search Markers
- Search Traits
- Search Mapped BACs

Sequences

- ESTs : Search
- Mapped BACs
- Genome
- Proteins
- Organelle

What's New in GDR?

- Unigene build 4 now available for **Malus, Fragaria, Rosa, Pyrus and Rubus**.
- New **local BLAST** server implemented.
- GDR redesigned. Any problems/comments? Please let us know!
- New **Rosaceae Community Forum** site.
- New **Comparative mapping and transferrable markers** site.
- New **RosCAP proposal** information portal available.
- Updated **peach physical map** available from **WebChrom** and **WebFPC**.
- GDR designed Nimblegen **Malus unigeneV4** array in production.
- Download the **Rosaceae/genus proteins**.
- New **Apple and Pear Polymorphism** data added.
- **Article on GDR** published in **Nucleic Acids Research**.
- New **Apricot Cmap, Apricot LxB 2007**.

Rosaceae Community News

- Funding Opportunities for Rosaceae in the **Specialty Crops Research Initiative**.
- **"Multiple Models for Rosaceae Genomics"** article by Shulaev *et al.* published in **Plant Physiology**.
- Download the **Rosaceae NRI 2008 Projects Reports**.
- Get the latest news on apple, peach and strawberry **whole genome sequencing**.
- Download the **minutes** and abstracts from the **4th International Rosaceae Genomics Conference** held at Chile in March 2008.
- **Plant Ontology Consortium** announced February 2008 release of **Plant Ontology** database.
- Download the **minutes** from the **USRosEXEC April 2008 Meeting**.
- Download the **minutes** from the **International Rosaceae Genomics Meeting** at the **2008 Plant and Animal Genome Conference**.
- Download the **minutes** from the **USRosEXEC Meeting** at the **2008 Plant and Animal Genome Conference**.
- **WSU and UW** announces **sequencing of a double haploid apple genome**.
- **Rosaceae standardized phenotyping** effort progress.

And there's more

The homepage has a number of important features.

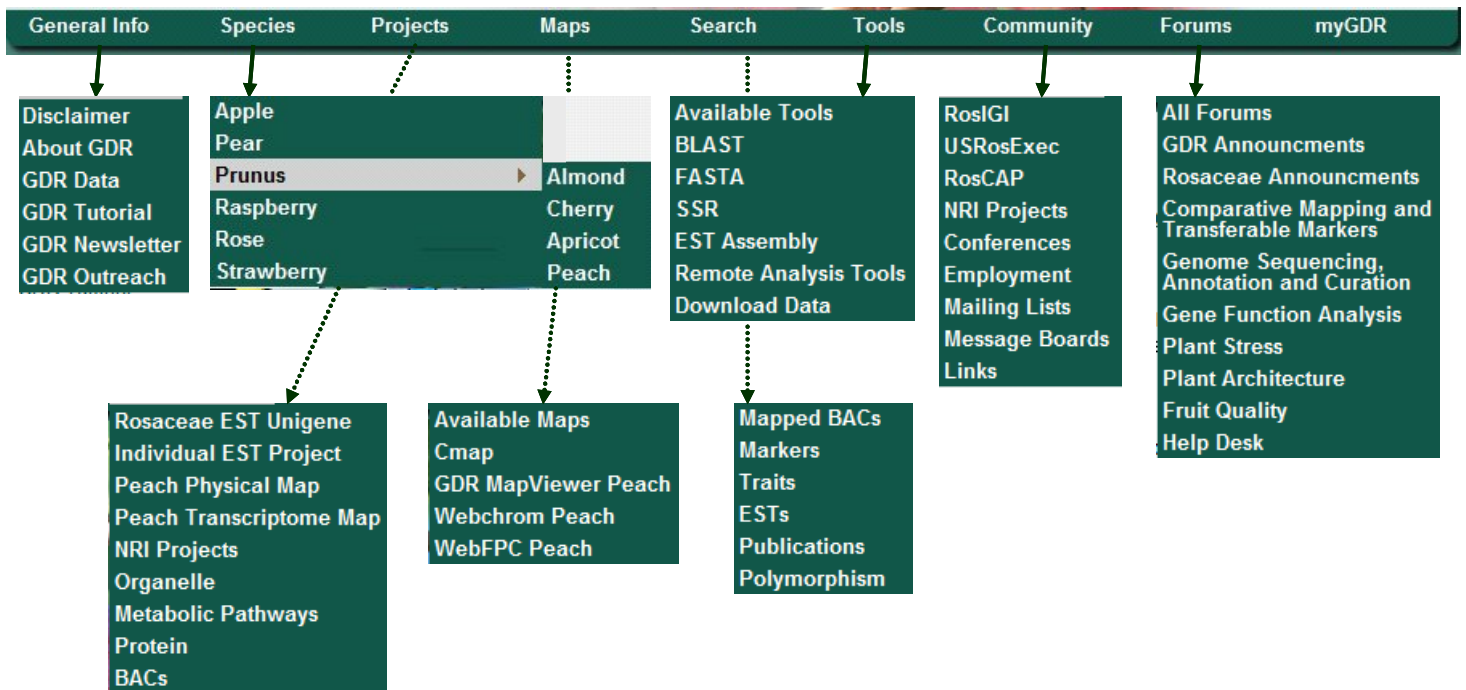
What's New in GDR lists new features of the database and is regularly updated.

Rosaceae Community News lists upcoming conferences, job opportunity, and various documents for the community.

The dark green bar and white links found at the top of the page are navigational tools. Similar links are expanded and listed for easy access along the light green bar at the left of the page.

The GDR provides the same navigation bar at the top of all of its pages to allow users to easily move between sections of the database.

The eight main sections, general info, species, projects, maps, search, tools, community and forums, each have a drop down menu with further options. To see this menu, move your pointer over the title. The last section, myGDR, will be developed in the future.



The tutorials provided systematically explore each of the sections available through this main menu, but are able to be done individually as well.

2. Please continue by clicking on the “About GDR” link under “general info” in the dark green navigation bar.

About GDR

The Genome Database for Rosaceae (GDR) is a curated and integrated web-based relational database. GDR contains comprehensive data of the genetically anchored peach physical map, annotated EST databases of apple, peach, almond, cherry, rose, raspberry and strawberry, Rosaceae maps and markers and all publicly available Rosaceae sequences. Annotations of ESTs include contig assembly, putative function, simple sequence repeats, ORFs, Gene Ontology and anchored position to the peach physical map where applicable. Our integrated map viewer provides graphical interface to the genetic, transcriptome and physical mapping information. We continue to add Rosaceae map data to CMap, a web-based tool that allows users to view comparisons of genetic and physical maps. ESTs, BACs and markers can be queried by various categories and the search result sites are linked to the integrated map viewer or to the WebFPC physical map sites. In addition to browsing and querying the database, users can compare their sequences with the annotated GDR sequences via a dedicated sequence similarity server running either the BLAST or FASTA algorithm, search their sequences for microsatellites using the SSR server or assemble their ESTs using the CAP3 Server.

Funding source: GDR has been financed by the NSF PGRP - Award #0320544.

Contact Us: If you have any problem, please contact us either by email or through our feedback form, we welcome your comments.

GDR Usage: In 2007, GDR was accessed by researchers from 44 countries with over 3.9 million pages accessed.

How to reference GDR:

- Sook Jung , Margaret Staton , Taein Lee, Anna Blenda, Randall Svancara, Albert Abbott , and Dorrie Main. GDR (Genome Database for Rosaceae): integrated web-database for Rosaceae genomics and genetics data. *Nucleic Acids Research*, 2008 January; 36(Database issue): D1034–D1040.
- Sook Jung , Christopher Jesudurai , Margaret Staton , Zhidian Du , Stephen Ficklin , Ilhyung Cho , Albert Abbott , Jeffrey Tomkins and Dorrie Main. 2004. GDR (Genome Database for Rosaceae): integrated web resources for Rosaceae genomics and genetics research. *BMC Bioinformatics* 2004, 5:130.

Other GDR publications:

- S. Jung, D. Main, M. Staton, I. Cho, T. Zhebentyayeva, P. Arus, A. Abbott. Synteny conservation between the prunus genome and both the present and ancestral Arabidopsis genomes. *BMC Genomics* 2006 7(1):8.
- Kevin M Folta , Margaret Staton, Philip J Stewart, Sook Jung , Dawn H Bies, Christopher Jesudurai and Dorrie Main. 2005. Expressed sequence tags (ESTs) and simple sequence repeat (SSR) markers from octoploid strawberry (*Fragaria × ananassa*). *BMC Plant Biology* 2005, 5:12
- Horn R, Lecouls AC, Callahan A, Dandekar A, Garay L, McCord P, Howad W, Chan H, Verde I, Main D, Jung S, Georgi L, Forrest S, Mook J, Zhebentyayeva T, Yu Y, Kim HR, Jesudurai C, Sosinski B, Arus P, Baird V, Parfitt D, Reighard G, Soorza R, Tomkins J, Wing R, Abbott AG. 2005. Candidate gene database and transcript map for peach, a model species for fruit trees. *Theor Appl Genet.* 2005 May;110(8):1419-28.
- Jung S, Abbott A, Jesudurai C, Tomkins J, Main D. 2005. Frequency, type, distribution and annotation of simple sequence repeats in Rosaceae ESTs. *Funct Integr Genomics.* 2005 Jul;5(3):138-43.

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This page provides a basic description of the GDR project and what types of resources it makes available to Rosaceae family researchers.

The funding sources are listed and links to the NSF award site.

GDR publication is also listed and is available online at the Bioinformatics site. This is another good resource for learning more about the GDR.

This page has hyperlinks to email or feedback forum. You can contact us either way. The GDR team welcomes any and all questions relating to the website or Rosaceae research.

3. Please continue by clicking on the “GDR Data” link under “general info” in the dark green navigation bar.

Data Overview

Rosaceae Mapping Data:

- Genetically Anchored Peach Physical Map:**
 The latest [peach physical map](#) (Apr 2008) contains 18387 BACs and 1367 contigs. A total of 149 contigs are anchored to the TxE map.
- Peach Transcriptome Map:**
 A total of 367 peach unigenes are anchored to genetic maps by hybridization to genetically anchored BACs. Of these, 216 unigenes are anchored to the TxE map. An additional 382 peach unigenes are anchored to genetic maps by hybridization to BACs that belong to genetically anchored BAC contigs.
- Rosaceae Genetic Maps:** Rosaceae genetic maps are available through CMap, the comparative map viewer. A total of over 1000 molecular markers and traits from various Rosaceae genetic maps and diversity studies are available from the [marker or trait search page](#).
- Data available for the three peach BACs that are completely sequenced.**
 Anchored ESTs/markers and the sequences for the three BACs (082118, 028F08 and pPn31C7) can be accessed from [BAC search site](#). An online annotation pipeline is currently under development by the GDR team.

Rosaceae EST Data:

- SNP Analysis:** The publicly available autoSNP software was used to analyze the most recent versions of the Rosaceae unigene and the individual genera unigenes. A summary of the results and downloadable data can be found [here](#).
- EST Assemblies:** Assemblies of publicly available ESTs are available for the family Rosaceae and the genera *Malus*, *Prunus*, *Fragaria*, *Rosa*, *Pyrus*, and *Rubus*. Homology data for each assembly with SWISS-PROT is available to browse and download. Homology with the NCBI nr protein database and other rosaceae ESTs will be available soon.
- EST Analysis:** Full bioinformatics analysis, including quality trimming, assembly, homology searching, microsatellite discovery and putative unigene development, is currently available for [Clemson Peach](#) and [Almond](#), [UF Octoploid Strawberry](#), and [CRA ISF Peach](#) ESTs. [Rose](#) and [blackberry](#) projects will be available soon.

Organism	Project	EST Number
Rosaceae	Rosaceae Assembly	359001
Fragaria	Fragaria Assembly	49132
<i>x ananassa</i> (octoploid strawberry)		
whole plant:	Kevin Folta - University of Florida	1505
<i>vesca</i> (diploid strawberry)		
unopened flower buds:	Tom Davis - University of New Hampshire	2717
Malus	Malus Assembly	249195
Prunus	Prunus Assembly	89166
<i>amygdalus</i> (almond)		
developing seed:	Albert Abbott - Clemson University	2794
<i>persica</i> (peach)		
fruit mesocarp:	Albert Abbott - Clemson University	9984
root:	Albert Abbott - Clemson University	Available Soon
shoot:	Albert Abbott - Clemson University	7085
fruit mesocarp:	Elisa Vendramin - CRA ISF	1667

This page provides an overview of some of the projects being done by the GDR bioinformatics team and other collaborating researchers.

The grey chart displays the different EST projects that are available through the GDR. Clicking on an EST project will take you to the main page for that project.

The Rosaceae Mapping Data in available in GDR are listed and the appropriate links are provided. The mapping section of the database is also linked through the main navigation bar and has its own tutorial, “Using GDR Mapping Resources.”

4. Please continue by clicking on the “Apple” link under “species” in the dark green navigation bar.

Each species page lists of and provides links to all the data for the species available in GDR, such as mapping, BAC, marker, and sequence data. Species page also provides links to remote sites for funded projects and other related links for the species.

5. Please continue by clicking on the “Links” link under “Community” in the dark green navigation bar.

This page is good resource for finding information not available on the GDR website. The links are broken into three main categories, plant genomics and/or germplasm databases, genetics and genomics institutions, and journals.