Abstract. MicroRNAs (miRNAs/miRs) are a family of small, endogenous and evolutionarily-conserved non-coding RNAs that are involved in the regulation of several cellular and functional processes. miRNAs can act as oncogenes or tumor suppressors in all types of cancer, and could be used as prognostic and diagnostic biomarkers. Databases and computational algorithms are behind the majority of the research performed on miRNAs. These tools assemble and curate the relevant information on miRNAs and present it in a user-friendly manner. The current review presents 14 online databases that address every aspect of miRNA cancer research. Certain databases focus on miRNAs and a particular type of cancer, while others analyze the behavior of miRNAs in different malignancies at the same time. Additional databases allow researchers to search for mutations in miRNAs or their targets, and to review the naming history of a particular miRNA. All these databases are open-access, and are a valuable tool for those researchers working with these molecules, particularly those who lack access to an advanced computational infrastructure.

1. Introduction
Since the finding that small non-coding RNAs in Caenorhabditis elegans repress gene expression by binding to mRNA after transcription, our view on gene regulation has changed. These important small regulatory RNAs are known as microRNAs (miRNAs/miRs) (1,2).

miRNA-mediated gene regulation involves 3 steps: i) Processing the transcripts into ~22-nucleotide long RNAs; ii) reaching the target mRNA guided by argonaute (Ago) proteins; and iii) repressing expression by inhibiting or degrading the target mRNA. In the animal kingdom, each species has a number of miRNA genes (3) that control gene expression in the majority of developmental and physiological processes (4).

Among the processes with miRNA-regulated gene expression are the immune response, signal transduction, and cell proliferation and differentiation, as well as other metabolic processes, including fat metabolism and insulin secretion (5). miRNAs mostly affect gene expression by degrading or repressing the target mRNA, although there is evidence that miRNAs can also act as enhancers of gene expression (6). The level of complementarity between mRNA and miRNA determines whether the target gene will be degraded or inhibited; near-perfect complementarity means that the mRNA will be degraded, while imperfect pairing and bulges mean that the mRNA will be inhibited (7). The pathways that miRNAs use to degrade mRNAs are well understood, as they behave similarly to small interfering RNAs (8).

Given their significant role in normal developmental and physiological processes, it follows that miRNAs also play an important role in cancer, either as oncogenes (oncomirs) or tumor suppressor genes. The deregulated expression of miRNA in human cancers has been observed since 2002, first in hematopoietic cancer (9) and later in solid tumors such as...
cancers, and colonic and rectal adenomas (10), as well as in malignant brain tumors (11), thyroid (12), breast (13) and lung (14,15) cancer, and hepatocellular carcinomas (16). miRNA expression profiles can be more informative and accurate for identifying certain types of cancer than the currently used methods (17), and miRNA expression profiles can aid in cancer classification and prognosis.

The growing body of transcriptomic data on miRNA expression has fueled the growth of resources that analyze miRNA expression patterns in normal tissues. Certain databases have been created using computational predictions and experimental results, for example miRBase (18), miRNAmap2.0 (19), mirGen (20), miRgator v2.0 (21) and miRecords (22). Other databases, such as TargetScan (2), PicTar (23) and TargetMiner (24), are based on algorithms designed to predict microRNA targets according to the complementary pairing with the target. Another database, TarBase (25), uses validated microRNA targets, while RNAhybrid (26) is based on hybridization between miRNA and mRNA. There are also databases that provide information on various diseases, such as MicroCosm (27), PhenomiR 2.0 (28) and mir2Disease (29). However, few databases are devoted to cancer-related miRNAs. The current review presents several recently published web-based tools for studying the association between miRNA and cancer. These databases are summarized in Table I.

2. SomamiR DB 2.0

Circular RNAs (circRNAs) and long non-coding RNA (lncRNA) can occasionally share target miRNAs with miRNAs. When this occurs, these molecules can form a competing endogenous RNA (ceRNA) network. There appears to be an association between the CNRNA network and cancer. As miRNA recognition depends on the sequence, it follows that mutations in either miRNAs or targets will affect the miRNA-ceRNA union (30).

SomamiR DB 2.0 (http://compbio.uthsc.edu/SomamiR/) was created to improve on the search of somatic mutations in miRNAs and ceRNAs (lncRNAs, circRNAs and miRNAs). The database searches for somatic mutations associated with cancer and allows researchers to perform a functional search of these mutations.

The database can be searched for somatic mutations in miRNA sequences, and offers the option of using miR2GO. miR2GO can perform a functional analysis of the importance of these mutations in miRNAs. Additionally, miR2GO offers miRmut2GO, an option that checks whether mutations on the miRNA produce changes in the target, and miRpair2GO, which offers a gene ontology analysis for the target gene sets of different miRNAs. SomamiR 2.0 can also search for miRNA mutations that have been experimentally identified by the crosslinking, ligation and sequencing of hybrids. The database can also search for mutations that have been experimentally detected using photoactivatable-ribonucleoside-enhanced crosslinking and immunoprecipitation, and high-throughput sequencing of RNA isolated by crosslinking immunoprecipitation.

Researchers can also search SomamiR 2.0 for predicted mutations on the 3′-untranslated regions (3′UTRs) of target genes, as well as lncRNAs and circRNAs that create or destroy putative target sites on miRNAs. The search for biological pathways affected by mutations in miRNA target sites shows cancer genes associated with Kyoto Encyclopedia of Genes and Genomes pathways. Finally, the database can be searched for mutations on miRNAs associated with cancer, or their target sites.

SomamiR 2.0 can be used to search for somatic and germ-line mutations in miRNAs or targets that are closely related to cancer if the genes were associated with cancer risk using GWAS (Genome-wide association study) or CGAS (candidate-gene association studies) and if there is experimental evidence that the mutation alters the miRNA functions. The mutations that comply with these criteria can be accessed by clicking on experimental evidence linking miRNA related polymorphisms with cancer or experimental evidence linking polymorphisms in miRNA with cancer, respectively. SomamiR is a tool that facilitates the search for cancer-related mutations and can be very a valuable tool in the understanding of this disease (31,32).

3. miRandola

Extracellular vesicles (exosomes, microvesicles and other membranous vesicles) can be found in a number of bodily fluids, including blood, urine and saliva, and are involved in cancer biology due to the role that they play in intercellular communication. Extracellular vesicles contain a variety of molecules, among which are miRNAs, lncRNAs, mRNAs and proteins, and these molecules are associated with the appearance of cancer, from the start of the tumor to angiogenesis, immunologic vigilance, drug resistance, invasion and metastasis (33).

miRandola (http://mirandola.ist.cnr.it) offers information on the miRNAs present in extracellular vesicles. Extracellular miRNAs are classified into four groups: miRNA-Ago2, miRNA-high-density lipoprotein (HDL), miRNA-exosome and miRNA-circulating. In the latter category are all those miRNAs that cannot be classified in the other 3 categories, as the study did not specify it.

To date, miRandola contains information from 271 studies, yielding 2,681 entries for non-coding RNAs (ncRNAs), among which are 174 miRNAs associated with Ago2, 1,618 circulating miRNAs, 868 miRNAs in exosomes, 21 miRNAs in HDL and 9 miRNAs in microvesicles. The database has 2,179 entries for 12 different types of body fluids: 1,096 for plasma, 983 for serum, 28 for blood, 15 for saliva, 6 for amniotic fluid, 7 for breast milk, 5 for cerebrospinal fluid, 6 for colostrum, 5 for peritoneal fluid, 4 for pleural effusions, 6 for seminal fluid and 18 for urine.

The database has fast and advanced search options. In the fast search, users can search by mature miRNA, ncRNA, miRNA family, type or RNA (Ago, circulating, exosome and HDL), disease, malignant cell line and potential biomarkers. The results can be exported as .xls or .csv. In the advanced search, users can select between miRNA families or mature miRNAs and sample type. The results for this database include links to the miRô database (34), which offers information on the associations between the miRNA and diseases, function, processes and tissues.
<table>
<thead>
<tr>
<th>Web based tool</th>
<th>Website</th>
<th>Last update</th>
<th>Usefulness</th>
<th>(Ref.)</th>
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<tr>
<td>SomamiR DB 2.0</td>
<td><a href="http://compbio.uthsc.edu/SomamiR/">http://compbio.uthsc.edu/SomamiR/</a></td>
<td>2016(^a)</td>
<td>Browse somatic mutations in:</td>
<td>(31)</td>
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<td></td>
<td></td>
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<td>-miRNA</td>
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<td></td>
<td>-Experimentally identified miRNA target sites by CLASH, PAR-CLIP, and HITS-CLIP experiments</td>
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<td></td>
<td></td>
<td></td>
<td>-Predicted miRNA target sites</td>
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<td>-Biological pathways</td>
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<td></td>
<td></td>
<td></td>
<td>-Genes associated with cancer risk that contain miRNA</td>
<td></td>
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<tr>
<td>miRandola</td>
<td><a href="http://mirandola.iit.cnr.it">http://mirandola.iit.cnr.it</a></td>
<td>June, 2015</td>
<td>Retrieve information on extracellular/circulating microRNAs, including:</td>
<td>(35)</td>
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<td></td>
<td></td>
<td></td>
<td>-mature miRNA</td>
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<td>-long non-coding RNA</td>
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<td>-miRNA family</td>
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<td>-miRNA type</td>
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<td>-malignant cell lines</td>
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<td></td>
<td></td>
<td>-potential biomarkers</td>
<td></td>
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<tr>
<td>PROGmiR</td>
<td><a href="http://www.compbio.iupui.edu/progmir">www.compbio.iupui.edu/progmir</a></td>
<td>December, 2012</td>
<td>Study the prognosis biomarker potential of different miRNAs in several types of cancer and overall survival query</td>
<td>(39)</td>
</tr>
<tr>
<td>AutomiRDB</td>
<td><a href="http://www.chen-lab.com/index.php">http://www.chen-lab.com/index.php</a></td>
<td>2014(^a)</td>
<td>Groups all the miRNAs that are experimentally confirmed to have an association with autophagy in cancer</td>
<td>(42)</td>
</tr>
<tr>
<td>OncomiRdB</td>
<td><a href="http://tdb.ccmrb.res.in/OncomiRdB/index.htm">http://tdb.ccmrb.res.in/OncomiRdB/index.htm</a></td>
<td>2014(^a)</td>
<td>Retrieve microRNAs with known breast cancer associations from miRNA databases, including miRBase, miR2Disease and PhenomiR</td>
<td>(43)</td>
</tr>
<tr>
<td>miRCancer</td>
<td><a href="http://mircancer.ecu.edu">http://mircancer.ecu.edu</a></td>
<td>December, 2015</td>
<td>Stores records of miRNA and cancer associations collected through data mining</td>
<td>(45)</td>
</tr>
<tr>
<td>CancerNet</td>
<td><a href="http://bis.zju.edu.cn/CancerNet">http://bis.zju.edu.cn/CancerNet</a></td>
<td>2015(^a)</td>
<td>Focuses on protein-protein interactions associated with cancer</td>
<td>(48)</td>
</tr>
<tr>
<td>canEvolve</td>
<td><a href="http://www.canevolve.org">http://www.canevolve.org</a></td>
<td>2013(^a)</td>
<td>Tool for oncogenicomic analysis</td>
<td>(51)</td>
</tr>
<tr>
<td>HNOCDB</td>
<td><a href="http://gyanxet.com/hno.html">http://gyanxet.com/hno.html</a></td>
<td>2012(^a)</td>
<td>Presents a classification of miRNAs, genes, and chromosomes involved in HNOC</td>
<td>(53)</td>
</tr>
<tr>
<td>miREC</td>
<td><a href="http://www.mirecdb.org">http://www.mirecdb.org</a></td>
<td>May, 2014</td>
<td>Search known miRNAs involved in endometrial cancer</td>
<td>(57)</td>
</tr>
<tr>
<td>Renal Cancer Gene DB</td>
<td><a href="http://www.juit.ac.in/attachments/jst/rcdb/homenew.html">http://www.juit.ac.in/attachments/jst/rcdb/homenew.html</a></td>
<td>2012(^a)</td>
<td>Contains miRNAs contributing to the etiology and pathogenesis of different types of renal cancer</td>
<td>(60)</td>
</tr>
<tr>
<td>Pancreatic Cancer DB</td>
<td><a href="http://www.pancreaticcancerdatabase.org">http://www.pancreaticcancerdatabase.org</a></td>
<td>July, 2015</td>
<td>Retrieve experimentally demonstrated molecular alterations of miRNA, protein and mRNA levels associated with pancreatic cancer</td>
<td>(63)</td>
</tr>
<tr>
<td>Sarcoma microRNA Expression DB</td>
<td><a href="http://www.oncomir.umn.edu/">http://www.oncomir.umn.edu/</a></td>
<td>2010(^a)</td>
<td>Presents differentially-expressed miRNAs in sarcoma</td>
<td>(65)</td>
</tr>
<tr>
<td>miRBase Tracker</td>
<td><a href="http://www.mirbasetracker.org">http://www.mirbasetracker.org</a></td>
<td>2014(^a)</td>
<td>Presents all the known miRNA sequences and variants, and keeps track of annotation changes for each miRNA entry</td>
<td>(66)</td>
</tr>
</tbody>
</table>

\(^a\)Publication date with no update on the website. DB, database; miR/miRNA, microRNA; CLASH, crosslinking, ligation, and sequencing of hybrids; PAR-CLIP, photoactivatable-ribonucleoside-enhanced crosslinking and immunoprecipitation; HITS-CLIP, high-throughput sequencing of RNA isolated by crosslinking immunoprecipitation; HNOC, head and neck and oral carcinoma.
miRandola has two notable tools: miRNAexpress, which makes a systematic comparison of the expression profiles of cellular and extracellular miRNAs, and miMETA, a tool for the meta-analysis of the miRandola database. miRandola specializes in circulating miRNAs, which are increasingly important as non-invasive biomarkers. The database is curated manually, and the data is recovered from the full paper, not just the abstract; this allows miRandola to offer more detailed information, such as method of analysis and a description of the experiment. Finally, the database offers links to other useful databases, including miRBase (www.mirbase.org), miRò (ferrolab.dmi.unict.it/miro/), ExoCarta (www.exocarta.org), and PubMed (www.ncbi.nlm.nih.gov/pubmed) (35,36).

4. PROGmiR

PROGmiR (www.compbio.iupui.edu/progmir) allows researchers to study the prognosis biomarker potential of various miRNAs in several types of cancer and overall survival (37). Information from Gene Expression Omnibus and The Cancer Genome Atlas is used, together with information on 16 different types of cancer to create survival Kaplan-Meier plots (38). The tool, which is intended for hypothesis-building only, can offer researchers information regarding the role that a certain miRNA plays in a certain cancer, whether it acts as an oncogene or a tumor suppressor, or both. PROGmiR is freely available online for non-commercial use. The database can be searched by miRNA ID, and then the users can select the cancer type for which they want to see the prognosis, or the 3- or 5-year survival rate. Each plot is created from one database, so as to avoid bias or other issues. Currently, the database has the profiles of 1,050 miRNAs from 3,117 samples from 16 types of cancer (39).

5. AutomiRDB

Autophagy is a cellular process that can lead to cell death, where a cell destroys proteins and other substances present in its cytoplasm. Autophagy can prevent cells from becoming cancerous, but it can also protect cancer cells from anticancer drugs (40). There is evidence that miRNAs have a significant role in autophagy regulation pathways, and this growing research field can aid in the development of novel therapies against cancer (41).

AutomiRDB (http://www.chen-lab.com/index.php) groups all the miRNAs that are experimentally confirmed to have an association with autophagy in cancer (42). The information in the database allows users to associate human miRNAs with target genes or proteins in different malignancies. To develop AutomiRDB, the published literature was searched for genes involved in autophagy in cancer, and predictive analysis was performed to determine which miRNAs would target those genes. The database includes 493 miRNAs associated with 90 genes or proteins associated with autophagy in 18 different types of cancer. The database is simple to use; the user only has to type the name of the miRNA in the query. The results page contains hyperlinks that lead to more information in other databases, including OMIM (http://www.omim.org) and UNIPROT (http://www.uniprot.org), among others (42).

6. OncomiRdbB

OncomiRdbB (tdb.ccmbr.res.in/OncomiRdbB/index.htm) (43) was built from 782 human and 246 mouse microRNAs that possessed known associations with breast cancer, which were retrieved from miRNA databases, including miRBase (18), miR2Disease (29) and PhenomiR 2.0 (28). The findings were validated using Taqman low density arrays that consisted of 667 human miRNAs and LNA™ arrays using human breast cancer samples. These tests yielded ~400 breast cancer miRNAs, as well as 34 miRNA sequences that had not been classified as such. To find the targets of the miRNAs involved in breast cancer, the developers compared the 3'UTR, 5'UTR and exonic regions of genes from different pathways using miRanda. This analysis revealed 711 potential target genes for humans and 490 for mice, which is more than can be found in other databases.

OncomiRdbB is a freely available web interface. Users can search miRNAs by name, accession number, target gene or sequence. The genes are classified according to their signaling pathway, and the database includes information on gene location within the chromosome, and alignment between the miRNA and the target, as well as links to the parental databases where the genes were retrieved (43).

7. miRCancer

miRCancer (http://mircancer.ecu.edu) is a database that stores records of miRNA and cancer associations collected through data mining (44). A rule-based approach was devised to analyze the title and abstract of 26,414 publications and to find full sentences or phrases that included the names of the miRNA and the cancer type, and any expression terms. The results of this data mining process were then corroborated by hand. miRCancer has records of >3,764 miRNA-cancer associations from 2,611 publications, which amounts to 236 miRNA expression profiles from 176 human cancers.

The rules used in text mining for this database resulted in a recall rate of 78%. One of the reasons for certain studies not being recalled can be attributed to the manner in which the study was written. The method used requires that the miRNA-cancer association be stated clearly, preferably in one sentence.

miRCancer is freely accessible online, and the database can be searched by miRNA name or cancer type, or a combination of both (45).

8. CancerNet

The biological processes that keep a cell in homeostasis result from a number of interactions between proteins. The alteration of these interactions can lead to pathological processes such as cancer (46).

There are several databases that deal with protein interactions, but CancerNet (http://bis.zju.edu.cn/CancerNet) is the first database to focus on the protein-protein interactions associated with cancer. As miRNAs have an important role in the mechanisms that regulate protein expression, building a database than integrates the miRNA regulation network with regard to specific cancer types is an important contribution.
miRNAs can act in synergy, i.e., several miRNAs can act on a single molecule, and this synergy can be different depending on the type of cancer (47). CancerNet takes this synergy into consideration and provides molecular interactions for 33 cancer types. The synergistic interactions included are 185,589 protein-protein interactions, 3,249,385 miRNA-target gene interactions and miRNA-miRNA interactions.

The CancerNet platform is simple to use and the queries can be made by molecule name and type of cancer, or by entering two molecules to find whether they have synergistic interactions. The results show a list with detailed information on these interactions, expression levels, functional similarity scores and the specificity of each interaction. The results can be viewed as a graph. Another function, the ‘GO Enrichment Analysis’ allows users to investigate the biological function of the genes that are of interest to them.

Applications for CancerNet include the study of specific interactions in cancer, and the study of the interactions between poorly annotated miRNAs and protein-coding genes (48).

9. canEvolve

Researchers working with cancer have benefited from the advances in the field of next-generation sequencing technology. There has been a massive amount of information derived from the genomic studies of healthy people and patients with certain diseases. Researchers searching for differences in the expression of miRNAs or genes in different types of cancer can occasionally struggle to find answers in the huge amount of data yielded by these technologies (49,50).

CanEvolve (http://www.canevolve.org) (51) is a database that can aid cancer researchers in answering their questions in a user-friendly manner by assisting them in searching through the data generated by massive sequencing experiments. The current version of canEvolve holds data from >18,000 patients in 127 datasets. The datasets include information such as gene expression, copy number alterations, miRNA expression, mutation, protein expression and protein-protein interactions. The database also has information on the differential expression of genes and miRNAs, changes in the number of copies, co-expression data, protein-protein interactions, metabolic and signaling pathways, and targets for transcription factors and miRNAs. The database also uses >200 algorithms, including LIMMA, dsChipSNP and ARACNE, to analyze different types of cancer.

CanEvolve can be searched by selecting the type of analysis and then the type of cancer and study. Next, users can enter genes or selected pathways, and click on ‘get results’. Results are shown as a heat map, graph or network depending on the desired analysis. The results can be downloaded as tables or R-data objects.

This database allows users to perform different types of integrative analysis and identify genes that are believed to induce or regulate tumorigenesis. It is also possible to perform a pathway meta-analysis from multiple studies of differentially-expressed genes and users are allowed to choose to review the enrichment of gene sets derived from MsigDB. CanEvolve is an excellent tool for oncogenic analysis and its tools will be extremely useful in the analysis of the huge amount of data that is being generated (51).

10. HNOCDDB

Head and neck and oral carcinomas (HNOCs) include oral, tongue, salivary gland, thyroid, pharyngeal, hypopharyngeal, nasopharyngeal, oropharyngeal and laryngeal cancer. These cancers are the sixth most common cancer in the world and are difficult to treat due to their location (52). Therefore, great efforts are being made to improve treatment and patient overlife.

HNOCDB (http://gyanxet.com/hno.html) (53) was created with information on genes and miRNAs involved in HNOC that have been validated. The genes were classified according to type of cancer, expression profile, miRNA ID and location in the chromosome. The database includes 451 genes and 109 miRNAs associated with HNOC.

Information can be accessed through 3 main links: ‘Genes’, ‘miRNAs’ and ‘chromosomes’, and then users can choose to see the association of the selected items with a certain type of cancer. In the chromosome section, users can choose the desired chromosome and then choose genes or miRNAs to study. In each section, the results are presented in a table that includes identifying data for the gene or miRNA, function and its alterations in this cancer.

HNOCDB is a database that presents a good classification of the miRNAs, genes and chromosomes involved in HNOC, and is a valuable tool for researchers that study this type of cancer (53).

11. miREC

Endometrial cancer (EC) is one of the most common gynecological cancers in developing countries (54), and as in all the other types of cancer that have been studied, EC has altered miRNA expression (55,56). Furthermore, as with all the other types of cancer, it is important to have a database that integrates all the information on miRNAs, gene annotation and expression, and the biomedical literature associated with this disease.

The miREC database (http://www.mirecdb.org) (57) has gathered information on all the known miRNAs involved in EC either by in silico prediction or experimental validation, which allows us to observe how miRNAs regulate genes in EC. The database (V 2.0) has information on 920 genes and 228 miRNAs. This information includes target genes and miRNAs, references to published scientific studies, literature citations for genes, miRNAs and gene-miRNA associations.

The database can be searched by gene or miRNA name, and has an advanced search option where users can introduce several parameters to refine their search. Results are presented in a table with links to other databases for more information. This database can also offer information on miRNA clusters through the identification of the functional correlation of associated miRNAs. This type of analysis is extremely useful, as it has been observed that miRNAs from the same cluster have common target genes, and that they can even be coexpressed and transcribed like a polycistron. There are databases that can do similar things, but miREC specializes in EC, and is a valuable tool for researchers looking for novel diagnostic tools and treatments for this disease (57).
12. Renal Cancer Gene Database

Renal cell carcinoma (RCC) is a common and often lethal urological cancer complicated by resistance to chemotherapy and radiotherapy (58). Although several molecular markers have been identified for RCC diagnosis and prognosis, it is not possible to predict tumor aggressiveness and metastasis (59).

To study renal cancer data, the Renal Cancer Gene Database integrated 240 protein-coding and 269 miRNAs associated with the pathogenesis of different RCC forms. Genes in this database are classified with regard to alteration in RCC, and deregulated miRNA is also included, with information associated with RCC type and metastatic or prognostic importance with common and unique miRNAs to RCC and other cancers.

The Renal Cancer Gene Database can be freely accessed (www.juit.ac.in/attachments/js/rcdb/homenew.html) (60) and uses a keyword search. Users can retrieve a list of genes with specific category and chromosomal location indexed by research studies and complemented with public databases such as Swissprot and Refseq, among others. In addition, users can perform comparative studies to find which genes are shared with other cancers and which genes are unique to RCC. This database is a good resource for researchers and physicians working in renal carcinoma (60).

13. Pancreatic Cancer Database

Pancreatic cancer is a leading cause of cancer-associated mortality worldwide (61). A wide range of alterations at the genome, transcriptome and proteome level have been reported for this malignancy. Although expression studies have proposed potential biomarkers for the diagnosis and prognosis of pancreatic cancer (62), there are no repositories and it is difficult to analyze a single gene in the different data and publications.

The Pancreatic Cancer Database provides an integrated web-based resource (www.pancreaticcancerdatabase.org) (63) containing data on the altered expression of 3,481 total genes in pancreatic cancer, from which 703 genes are altered at the mRNA and protein levels, 570 genes are altered at the protein level only and 1,982 genes are altered at the mRNA level only. In addition, the Pancreatic Cancer Database also contains 226 miRNAs associated with pancreatic cancer tissues and cell lines with an external link to miRBase. The Pancreatic Cancer Database was constructed using PHP (http://www.php.net) as a server and MySQL (http://www.mysql.com) as a storage system.

Users can quickly access the Pancreatic Cancer Database with genes, proteins, molecular alterations, type of cancer, different cell lines and experimental methods using the ‘browse’ option to navigate through the alterations described at the RNA, protein or miRNA levels (63).

14. Sarcoma microRNA Expression Database

Human sarcomas are rare, aggressive and invasive tumors that comprise a number of histological subtypes, with few markers for diagnostic and classification. miRNA expression patterns have been proposed as novel biomarkers for diagnosis, classification and prognosis in tumors, but are poorly understood in sarcomas (64).

The Sarcoma microRNA Expression Database (www.oncomir.umn.edu/) (65) was developed as a repository of human sarcoma miRNA expression patterns to address the lagoons in biological and bioinformatics knowledge. This database allows quick access to miRNA expression profiles of 310 tumor tissue samples that represent 22 different types of sarcomas.

The Sarcoma microRNA Expression Database presents the results as a heat map with text and numerical formats, and also provides statistical fold changes and P-values for differentially-expressed miRNAs. Since numerical values can be difficult to interpret, the heat map presents the option of color-coding to represent data in absolute and relative formats (65).

15. miRBase Tracker

miRBase (www.mirbasetracker.org) (66) is a database that includes all the known miRNA sequences and variants, and keeps track of annotation changes for each miRNA entry. Due to the large amount of research on miRNAs, the number of sequences and annotations has increased, from 218 precursor and 218 mature miRNAs from five different species in 2002, to 28,645 precursor and 35,828 mature miRNAs from 223 species in 2014.

The rapid growth in the number of miRNA sequences and annotation, as well as the identification of isomiRs resulted in a degree of confusion with miRNA identity. A proportion of this confusion comes from the use of outdated miRNA names in as much as 25% of the publications, as well as in ambiguities such as using the same name for different miRNA sequences.

The miRBase Tracker works with the miRBase database by keeping track of all the changes undergone by a miRNA sequence through different versions of miRBase in a MySQL base. Users can search the database by miRNA history to observe all the changes and annotations the sequence has gone through. An miRNA update search will yield the most recent annotation for the sequence, and a search by miRBase release compares precursor and mature miRNAs from any species between two miRBase releases, showing the changes undergone by the sequence (66).

16. Conclusions

This review shows a variety of user-friendly, open-access web-based tools that connect miRNAs with cancer through experimentally supported results. Although the majority of aspects of the mechanisms surrounding the interactions between miRNAs and cancer remain unknown, these databases can assist researchers to answer these questions. Databases focused on a particular type of cancer such as HNOCDB, miREC, the Renal Cancer Gene Database, the Pancreatic Cancer Database and the Sarcoma microRNA Expression Database are useful to those researchers who want to study the role of miRNAs in cancer biogenesis and evolution. Other databases (CanEvolve, CancerNet, miRCancer, OncomiRdbd, AutomiRdb and ProgmiR) that study the role of miRNAs in different tumors can be used to see the bigger picture of a particular miRNA in different pathologies.
miRandola and SomamiR are general databases that could play an important role in cancer research. MiRandola presents information on circulating miRNAs that may have an important role in metastasis and may be used to develop non-invasive biomarkers, while SomamiR allows researchers to find mutations that could lead to the development of malignancies.

The miRBase Tracker is not a cancer database, but it was included due to its efforts to unify miRNA naming conventions. This database allows those researchers working with miRNAs to track the name of a particular molecule through the years, thus increasing the scope of the research.

These web-based tools concerning miRNAs and cancer are a central resource for researchers; they are extremely useful in the validation of these targets and in the development of novel clinical cancer biomarkers, and ultimately, novel therapeutic interventions. More importantly, these databases assist researchers in navigating the enormous amounts of data available on miRNAs and cancer by providing curated, specific information in one place.

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