

# Towards a Standard Reference Database for Computer-aided Mammography

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## ABSTRACT

Because of the lack of mammography databases with a large amount of codified images and identified characteristics like pathology, type of breast tissue, and abnormality, there is a problem for the development of robust systems for computer-aided diagnosis. Integrated to the Image Retrieval in Medical Applications (IRMA) project, we present an available mammography database developed from the union of: The Mammographic Image Analysis Society Digital Mammogram Database (MIAS), The Digital Database for Screening Mammography (DDSM), the Lawrence Livermore National Laboratory (LLNL), and routine images from the Rheinisch-Westfälische Technische Hochschule (RWTH) Aachen. Using the IRMA code, standardized coding of tissue type, tumor staging, and lesion description was developed according to the American College of Radiology (ACR) tissue codes and the ACR breast imaging reporting and data system (BI-RADS). The import was done automatically using scripts for image download, file format conversion, file name, web page and information file browsing. Disregarding the resolution, this resulted in a total of 10,509 reference images, and 6,767 images are associated with an IRMA contour information feature file. In accordance to the respective license agreements, the database will be made freely available for research purposes, and may be used for image based evaluation campaigns such as the Cross Language Evaluation Forum (CLEF). We have also shown that it can be extended easily with further cases imported from a picture archiving and communication system (PACS).

**Keywords:** Database Construction, Mammography, Ground Truth, Computer-Aided Diagnosis

## 1. INTRODUCTION

In occidental countries, breast cancer represents one of the main causes of death among women [1]. Its symptoms are nodules or tumors in the breast (Brazil National Cancer Institute, <http://www.inca.gov.br>). The early detection is the most effective way to reduce mortality, and mammography is the best method of screening for breast cancer because it can show lesions in their initial phases. The diagnosis of these lesions is made by a radiologist and due to the fast and continuous advances in computer technology as well as the conversion to the digital format of mammographies there is an increasing interest in computer-aided diagnosis (CAD) systems [2].

An effective CAD system, i.e., a system that clearly identifies position, size, and staging of lesions like microcalcifications and masses in x-ray mammographies, must be evaluated using a large number of reference images with approved diagnostics (ground truth), e.g., [3].

However, current studies are based on rather small sets of data. For instance, Zwigelaar et al. used synthetic data and 15 mammographies of the Mammographic Image Analysis Society digital mammogram database (MIAS, <http://peipa.essex.ac.uk/ipa/pix/mias/>) to detect linear structures and classify them into the anatomical types: vessels, ducts, and spicules [4]. Christoyianni et al. detected the exact location of circumscribed masses in 22 images of MIAS

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Database	Resolution						File type	
	x-min	x-max	y-min	y-max	g-min	g-max	format	standard
DDSM	1,411	5,641	3,256	7,111	12 bit	12 bit	LJPEG	no
MIAS	334	1,000	802	1,024	8 bit	8 bit	PNG	yes
LLNL	700	4,494	2,828	6,874	12 bit	12 bit	ICS	no
RWTH	1,582	4,129	3,382	5,928	12 bit	12 bit	DICOM	yes

**Table 1:** Resolution and image type of databases

Database	Anatomy		Direction		Tissue type	Tumor staging	Lesion description	
	left	right	CC	ML			position	multiple
DDSM	4,918	4,915	4,917	4,916	ACR	yes	chain code	yes
MIAS	161	161	0	322	yes	yes	circle	yes
LLNL	93	91	92	92	yes	yes	chain code	yes
RWTH	81	85	85	79	ACR	BI-RADS	none	---

**Table 2:** Anatomy, direction, and biosystem information

database [5]. Strickland & Hahn employed wavelet decomposition in 40 mammographies of Nijmegen database to detect micro-calcifications [6]. However, this database is no longer available. Arodz et al. achieved pattern recognition methods such as adaptive boosting and support vector machines to detect micro-calcifications and masses in 168 abnormal mammographies taken from the Digital Database for Screening Mammography (DDSM, <http://marathon.csee.usf.edu/mammography/database.html>) [7]. Also based on DDSM, Eltonsy et al. tried to localize potentially malignant masses in a set of 540 images [8].

Although even large databases for mammography – such as DDSM with about 10,000 images – are public available, the problem for researchers needing reference data are manifold. Resolution, tissue classification and pathology description and/or localization of the region of interest (ROI) are not standardized, and may be partly erroneous. Resulting from different formats, databases cannot be merged, and for particular algorithms and CAD approaches, an insufficient number of cases may exist in each of the database, individually. Therefore, generation of classified mammography collections is still in the focus of recent research [9].

Regarding the large number of images, image handling and image data management becomes difficult, too. Researches must face the problem of selecting appropriate cases and presenting results of the computation visually in a transparent and user-friendly way. However, novel methodologies of visual data management are currently developed in the field of content-based image retrieval (CBIR) and image data mining [10]; for reviews on CBIR and medical CBIR cf. the publications of Smeulders et al. [11] and Müller et al. [12], respectively.

Based on the Image Retrieval in Medical Applications (IRMA, <http://irma-project.org>) framework [13], we aim at defining a unified database structure and coding scheme for mammographic radiographs associated with diagnostic information that can be filled consistently with images from the various databases available as well as mammography data from Picture Archiving and Communication Systems (PACS) as they are used in routine of hospitals and health centers. The unified reference database will be used for CAD system evaluation and other evaluation campaigns such as the automatic image annotation task in the Cross Language Evaluation Forum (CLEF, <http://www.clef-campaign.org>) [14, 15].

## 2. MATERIALS AND METHODS

In this section, we present details of the databases that we combine and explain how the images were imported into the IRMA system and unambiguously classified using the IRMA coding scheme.

### 2.1. Freely Accessible Mammography Databases

An overview of the data available for this study is given in Tables 1 and 2.

**DDSM.** The DDSM database [16] officially contains 2,479 studies (695 normal, 870 benign, and 914 cancerous cases). Each study includes two images of each breast, acquired in craniocaudal (CC) and mediolateral (ML) views that have been scanned from the film-based sources by four different scanners with a resolution between 50 and 42 microns. This results in total of 9,916 radiographs. These images are coded with an algorithm according to the lossless Joint Pictures Expert Group (JPEG) standard and had to be converted into a standard file format with special software that is provided in C source code at the DDSM web page.

For all cases, there are additional plain text files with information on the type of digitizer and a staging of the tissue density according to the American College of Radiology (ACR). Where appropriate, information about the lesions types and a chain code with the localization and delineation of these lesions are also provided.

**MIAS.** The MIAS database [17] is available only for researches purposes and contains 322 mammography images, all of them acquired in mediolateral view. Initially scanned from film with a resolution of 50 microns, all images were reduced to 200 microns and clipped / padded so that they fit into a 1,024 x 1,024 bounding box. The image files are available in the portable network graphics (PNG) format and annotated with the following details: a database reference number indicating left and right breast, character of background tissue, pathology, class of lesion present and coordinates as well as size of these lesions.

**LLNL.** The LLNL database [18] contains 197 mammography images (4 images per patient), all of them digitized at 35 microns per pixel. The images are stored in the image cytometry standard (ICS) format and had to be converted into a standard file format with a provided source code for a program that converts images in the ICS format to the portable grey map (PGM) format. For 190 images there is available a plain text file containing patient stats, biopsy results and ground truth comments.

**RWTH.** In order to evaluate the extensibility of mammogram reference resources, 170 cases were extracted arbitrarily from the PACS at the Department of Diagnostic Radiology, University Hospital, Aachen University of Technology (RWTH), Aachen, Germany. These images were acquired digitally using a General Electric Senographe operating with low beam energy about 26 to 32 kV and with a phosphor storage system from Fuji/Philips capable of recording 7 lp/mm. The cassette was read using a Philips PCR Eleva CosimaX. If available, a free text diagnosis in German describing the breast examination, pathology, type of tissue and lesion was included along with the digital imaging and communications in medicine (DICOM) files.

**OTHERS.** There are only few other databases that are public available and have been used for research. In some researches, the Nijmegen Database was used (e.g. by Heinlein et al. [19]), but since March 2000 it isn't available anymore. Others databases, e.g., the massive database provided by National Digital Medical Archive (NMDA) that holds over a million mammography images [20], are not freely available, and might be included in a second step.

## 2.2. The IRMA System

The IRMA project aims at developing and implementing high-level methods for content-based image retrieval (CBIR) with prototypal application to medico-diagnostic tasks on a radiological image archive [13]. Beyond the mammographies, there are currently more than 20,000 diagnostic images in the IRMA database, which are used for image retrieval [10,14].

In IRMA, all images are coded according to a mono-hierarchical, multi-axial coding scheme [21]. The four axes, each having three to four hierarchical positions, describe the

- *technique*: image modality,
- *direction*: body orientation,
- *anatomy*: body region examined, and the
- *biosystem*: biological system examined,

and result in a unique string of 13 digits: TTTT-DDD-AAA-BBB.

IRMA code		Tissue density	IRMA code		Tumor staging	IRMA	Type of lesion
xBB	ACR	description	BxB	BI-RADS	description	BBx	description
0	---	unspecified	0	BI-RADS 0	unspecified	0	unspecified
1-c	---	already in use	1	BI-RADS 1	normal	1	calcification, unspecified
d	ACR-1	fat transparent system	2	BI-RADS 2	benign	2	micro-calcification
e	ACR-2	fibroid glands system	3	BI-RADS 3	probably benign	3	macro-calcification
f	ACR-3	heterogeneously dense system	4	BI-RADS 4	suspiciously abnormal	4	circumscribed mass
g	ACR-4	extremely dense system	5	BI-RADS 5	malignant	5	spiculated mass
h	ACR-3/4	dense system				6	other mass
						7	architectural distortion
						8	asymmetry

**Table 3:** IRMA codes for the biosystem

**Technique.** The IRMA technical code for mammography is TTTT = 11xx, where 11 means x-ray, plain radiography, and the two remaining code positions are used to capture the nature of images (1 = directly digital, 2 = secondarily digitized) and their resolution (e.g., 42, 43.5, 50, or 200 microns).

**Direction.** According to the coding scheme, the directions for breast imaging, i.e., CC and ML, are denoted by DDD = 310 (axial – craniocaudal – unspecified) and DDD = 410 (other orientation – oblique – unspecified), respectively.

**Anatomy.** The anatomy axis of the IRMA code is used to differ left from right breast using the codes AAA = 610 (breast or mamma – right breast – unspecified) and AAA = 620 (breast or mamma – left breast – unspecified), respectively.

**Biosystem.** The biosystem axis was extended in order to capture tissue density, tumor staging, and lesion description (Table 3). The first position describes the tissue type according to the ACR classes. For instance, if the breast is almost entirely fat or rather scattered with fibro glandular densities, ACR-1 or ACR-2 is appropriate, respectively. The ACR-3/4 class (dense system) was defined to import the MIAS images. The second IRMA code position captures the tumor staging according to the ACR breast imaging reporting and data system (BI-RADS) [22, 23], where, for example, a breast that needs additional imaging evaluation is coded (BI-RADS 0). Finally, the third code position refers to type of lesion. According to the BI-RADS system, eight descriptions currently are defined (Table 3) which, according to the structure of the IRMA code, can be extended easily if demanded.

**Lesion Localization and Morphologic Description.** The IRMA framework provides a database that hosts images as well as their derived image features. For instance, texture signatures are associated with each image in order to allow fast CBIR access. Internally, this link is established based on the IRMA identifier, which is uniquely assigned to each database element. This relation is used to store one or more lesions descriptions, in form of

1. *circle*, described by its center coordinates and radius;
2. *contour points*, a list of (x,y)-coordinates;
3. *chain code*, a starting coordinate (x,y) followed by a sequence of numbers in [1..8] describing the direction to the adjacent contour point; or
4. *masking image*, a binary image with the same x-,y-dimensions as the mammography, where 0 and 1 denote “background” and “lesion”, respectively.

### 2.3. Integration of Databases

Integrating the images from DDSM, MIAS, LLNL and RWTH into the IRMA system, the IRMA code was set automatically, using the descriptions from the databases. The database-specific conversion problems are addressed in this chapter.

Code alteration	Description	Number
ACR 1 → ACR 2	fat transparent → fibroid glands	98
ACR 1 → ACR 3	fat transparent → heterogeneously dense	12
ACR 1 → ACR 4	fat transparent → extremely dense	1
ACR 2 → ACR 1	fibroid glands → fat transparent	102
ACR 2 → ACR 3	fibroid glands → heterogeneously dense	36
ACR 2 → ACR 4	fibroid glands → extremely dense	2
ACR 3 → ACR 1	heterogeneously dense → fat transparent	10
ACR 3 → ACR 2	heterogeneously dense → fibroid glands	136
ACR 3 → ACR 4	heterogeneously dense → extremely dense	4
ACR 4 → ACR 1	extremely dense → fat transparent	1
ACR 4 → ACR 2	extremely dense → fibroid glands	12
ACR 4 → ACR 3	extremely dense → heterogeneously dense	61
Total		475

**Table 4:** Correction of DDSM tissue classification

**DDSM.** The IRMA code was automatically generated based on bitmap file name as it contains anatomic and directional information, and the “.ics” file, which contains density information, and – if present – the “overlay” file, which stores diagnosis-related data, including the chain-code contour data for marking regions of interest (ROIs) with pathologies.

We encountered some problems with the data: first of all, 10 cases and 5 cases are missing from volumes benign\_01 and benign\_02, respectively. Furthermore, case 1658 from volume cancer\_11 was inaccessible for download. Therefore, 695 normal, 855 benign, and 913 malignant cases were available, with a total of 2,463 cases and 9,852 images. Processing the medical information yielded unproven pathologies for 15 images, which were excluded. In addition, case 1825 of volume cancer\_11 has density 0. Images with pathology “benign\_without\_feedback” were classified as “benign”.

The original format used to encode the images is LJPEG, and the authors also provide source code to extract the raw image data. Although the website states that each image uses 12bit quantization for grayscale intensities, 3,130 images violate the intensity boundaries of [0..4095]. Unfortunately, there are different patterns among these violations. Some images contain single outliers, while others fully utilize the extended value range. Therefore, an automatic conversion algorithm was applied, which first checks for outliers by inspecting a median-filtered version of the image. If only outliers caused the violation, these pixels are simply cut off. If a violation still exists, a check is performed if the image using the range fully. If the resulting image is still too dark (meaning that the found boundaries are still caused by surviving outliers), a grayscale stretching is performed based on a histogram which drops 1/64th of the pixels at the upper end. After the re-estimation of the upper grayscale boundary, the images were cropped and scaled to fit a 1024×1024 bounding box, and transferred into an 8bit grayscale intensity range.

Handling the data within the IRMA system, it became obvious that tissue classification partly was inconsistent and crosschecked by a trained radiologist. In total, ACR tissue code was altered by the physician for 475 images (Table 4).

**MIAS.** The codification was done manually according to the database description on the Internet. Initially the ACR tissue type was converted as follows:

- F (MIAS fatty) → d (ACR1);
- G (MIAS fatty-glandular) → e(ACR2);
- D (MIAS dense-glandular) → h (ACR3/4).

However, it turned out that this conversion does not result in correct ACR codes. From the 322 images, 187 had to be corrected by an experienced radiologist (Table 5). Furthermore, three images were corrected in BI-RADS classification from normal to benign with micro-calcifications.

**LLNL.** According to the information provided by text files, like direction of x-ray, breast anatomy, tissue type, pathology and lesion, the IRMA codes were associated. From the 190 images available, two images showing amputated

Code alteration	Description	Number
ACR 1 → ACR 2	fatty → fibroid glands	24
ACR 1 → ACR 3	fatty → heterogeneously dense	1
ACR 2 → ACR 3	fatty-glandular → heterogeneously dense	45
ACR 2 → ACR 4	fatty-glandular → extremely dense	4
ACR 3/4 → ACR 2	dense glandular → fibroid glands	5
ACR 3/4 → ACR 3	dense glandular → heterogeneously dense	38
ACR 3/4 → ACR 4	dense glandular → extremely dense	70
Total		187

**Table 5:** Correction of MIAS tissue classification

breasts and another four images with breasts containing silicone were excluded. Also, 2 images that had no diagnostic were inspected and codified manually by the trained radiologist.

**RWTH.** All the IRMA codes were manually extracted from the DICOM files. From the 170 available images, 54 were provided with full coding information, 81 had incomplete codes (tissue density or type of lesion was missing), and for 35 a diagnosis was not included. A physician inspected all these incomplete images and set the IRMA code.

### 3. RESULTS

In total 10,509 images (without differentiating the resolution) are now available for evaluation of mammography CAD systems. As it can be observed from Tables 6-8, the merged database holds examples for all categories and their combinations, which is not the case when restricting to only one data source.

Figures 1 and 2 exemplify the benefits of the new mammography reference database. All web-interfaces can be directly used to explore the database. For example, Fig. 1 shows a screen shot of the IRMA code statistics, a web-interface that lists all available code classes. Merging of groups can be easily computed using the wildcards in the parameter field [24]. Via the “view” buttons, this interface is linked to the IRMA code browser (Fig. 2), a web-interface for interactive exploration of the images. Nonetheless, the new resource is easily extensible, and further databases will be included as they become freely available for scientific research.

In total 10,509 images out of 323 code classes (disregarding the image resolution) are now available for evaluation of mammography CAD systems. As it can be observed from Tables 6-7, the merged database holds examples for all categories and their combinations, which is not the case when restricting to only one data source. As can be further deduced from Table 1, the majority of illustrations are still published in grayscale (87.86%). If multi-panel illustrations that contain at least one colored component were counted as if all components are colored, the number of grayscale panels is still above 80%. Similarly, the majority of illustrations are annotated with text, arrows, or other symbols which may cover image information and affect the textural feature extraction.

### 4. DISCUSSION

Although evaluation of medical image analysis should rely on large sets of ground truth data, research in computer-assisted mammography reading is often based on less than 30 images [4, 5]. Based on the IRMA framework, we defined a coding scheme according to the ACR standards and showed how existing image resources are integrated (DDSM, MIAS, LLNL and RWTH). More precisely, we provided a scheme to integrate available mammography databases using standardized description of imaging modality and resolution, orientation and view, left and right position of breast, tissue type, tumor staging and lesion description as well as lesion positions regardless whether these positions are coded by some boundary points, a bounding circle, or a complete chain code. Since the proposed scheme is based on international ACR and BI-RADS codes, it is extensible and can be used for future evaluation of CAD system. However, some problems arose from the fact that existing resources does not provide complete coding information, or hide this information in file names and additional description files.

Database	Nominal	IRMA code BBB = xBB					Converted	
		d	e	f	g	h	total	percent
DDSM	9,916	1,252	3,691	2,896	1,994	0	9,833	99.2
MIAS (before corrections)	322	80 (105)	84 (104)	84 (0)	74 (0)	0 (113)	322	100.0
LLNL	197	12	84	68	20	0	184	93.4
RWTH	170	48	78	42	2	0	170	100.0
IRMA	10,615	1,395	4,043	3,048	2,023	113	10,509	99.0

**Table 6:** Statistics of issue classes after integration

Database	IRMA code BBB = BxB						Converted
	0	1	2	3	4	5	total
DDSM	0	6,181	1,848	0	0	1,804	9,833
MIAS (before corrections)	0	206 (209)	64 (61)	0	0	52	322
LLNL	6	47	111	6	0	14	184
RWTH	2	69	87	6	6	0	170
IRMA	8	6,503	2,110	12	6	1,870	10,509

**Table 7:** Statistics of tumor staging after integration

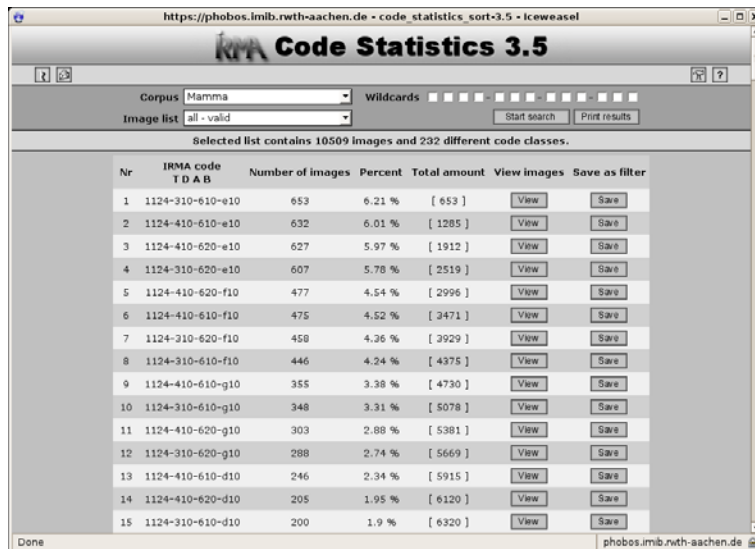
Database	IRMA code BBB = BBx									Converted
	0	1	2	3	4	5	6	7	8	total
DDSM	6,181	1,488	0	0	478	547	1,139	0	0	9,833
MIAS (before corrections)	206 (209)	23	0	3 (0)	23	19	14	19	15	322
LLNL	65	112	4	0	0	0	0	0	3	184
RWTH	71	0	40	43	4	2	0	10	0	170
IRMA	6,523	1,623	44	46	505	568	1,153	29	18	10,509

**Table 8:** Statistics of lesion types after integration

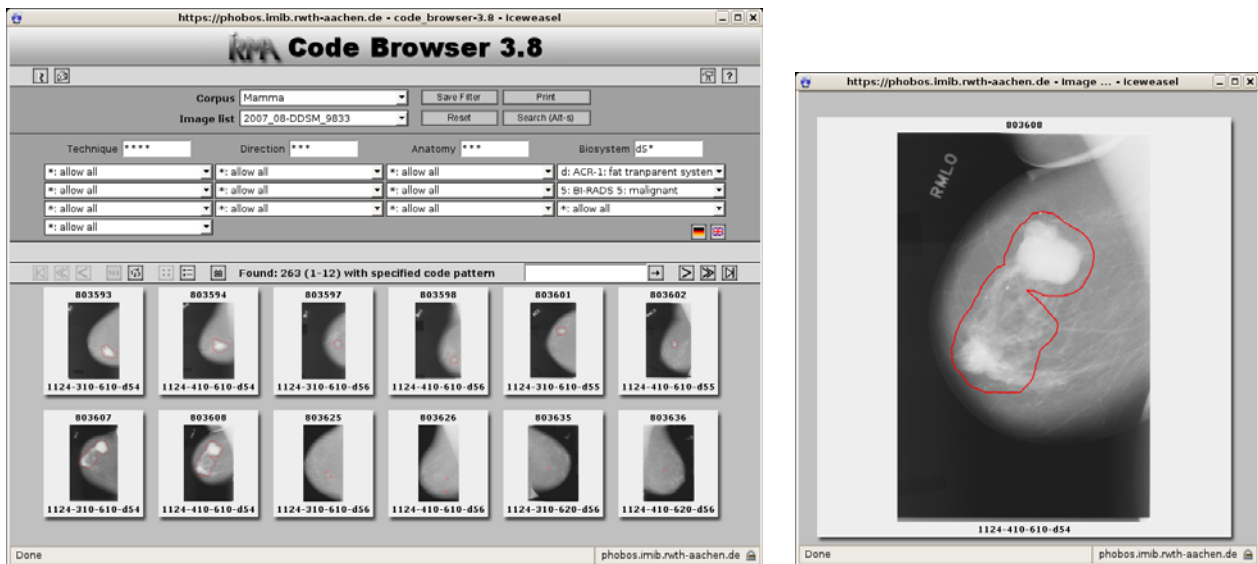
Building a mammography database with a large number of images is proposed by some researches [9], which intend also to provide information about the patients, like age, menstrual history and breast cancer occurrences in the family plus a ground truth data in XML format. With our database, containing until now 10,509 reference images, all the information needed by radiologists to their aid or for the implementation of retrieval and CAD systems is already given by the codified tissue density, lesion staging and type of lesions, without the need of extra toolboxes.

## 5. CONCLUSION

Based on international standards such as ACR and BI-RADS, we provided a scheme to integrate available mammography databases using standardized description of imaging modality and resolution, orientation and view, left and right position of breast, tissue type, tumor staging and lesion description as well as lesion positions regardless whether these positions are coded by some boundary points, a bounding circle, or a complete chain code. Integrating different resources that are freely available in the Internet, our database currently holds 10,509 images from 232 different code classes. Based on this unified database, researches in CAD systems can improve with the implementation of algorithms for tissue characterization.



**Figure 1:** IRMA Web-interface for code statistics. Currently, the corpus “Mamma” contains 10,509 images from 232 different code classes.



**Figure 2:** IRMA Web-interface for database browsing. Clicking on any icon in the overview (left) opens a separate window with more details (right). Diagnostic information is automatically overlaid.

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