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# Dereplication strategies in natural product research: How many tools and methodologies behind the same concept?

Jane Hubert<sup>a\*</sup>, Jean-Marc Nuzillard<sup>a</sup>, Jean-Hugues Renault<sup>a</sup>

#### **AUTHOR ADDRESS**

<sup>a</sup>Institut de Chimie Moléculaire de Reims (UMR CNRS 7312), SFR CAP'SANTE, UFR de

Pharmacie, Université de Reims Champagne-Ardenne, Reims, France

\*Corresponding author: jane.hubert@univ-reims.fr

#### ABSTRACT

The development of new drugs will certainly benefit from an ever improving knowledge of the living beings chemistry. However, identification of drugable molecules within the immense biodiversity of forests, soils or oceans still requires considerable investments in technical equipments, time and human resources. An important part of this process is the quick identification of known substances in order to concentrate the efforts on the discovery of new ones. A range of "dereplication" procedures are currently emerging to meet this challenge as key strategies to improve the performance of natural product screening programs. Initially defined in 1990 as "a process of quickly identifying known chemotypes", dereplication is today a not so univocal concept and has evolved over the last years in different ways. The present review covers all dereplication-related sudies in natural product research from 1990 to 2014. Its writing brought to light five distinct dereplication workflows that can be characterized by the nature of starting materials, by the key analytical technique, and above all by the final objective. Dereplication can be used as an untargeted workflow for the rapid identification of the major compounds whatever their chemical class in a single sample or for the acceleration of bioactivity-guided fractionation procedures. In other cases dereplication is fully integrated in metabolomic studies for the untargeted chemical profiling of natural extract collections or for the targeted identification of a predetermined class of metabolites. Finally a quite distinct dereplication approach mainly based on gene-sequence analyses is frequently used for the taxonomic identification of microbial strains.

#### **KEYWORDS**

Dereplication, natural products, metabolomics, drug discovery, taxonomic classification.

#### Introduction

Living beings provide a unique diversity of secondary metabolites that are still waiting to be investigated and exploited for the discovery of new drugs, cosmetic ingredients, nutraceutics or bio-sourced materials. The performance of natural product screening programs remains however strongly limited by the equipment, labor and time investments required for isolation and structural elucidation of novel metabolites with potent biological properties. Significant advances have been made over the last decades to improve the resolution and sensitivity of analytical methods for the study of small molecules in natural extracts. Very efficient hyphenation systems between separation and spectroscopic instruments now enable the identification of highly complex molecular structures even within complex mixtures of compounds. Despite these efforts, very serious issues, such as the frequent rediscovery of known compounds after time consuming purification and identification steps, the redundant investigation of taxonomically ambiguous microbial strains or the investigation of irrelevant crude materials regarding a selected biological application, still result in a considerable waste of time. Consequently, many pharmaceutical companies have considerably slowed down or even terminated their natural product research activities.

Nevertheless, the search for novel chemical structures within natural resources continues to be a crucial strategy for the discovery of novel therapeutic drugs. Moreover, remembering that more than 300 000 secondary metabolites have been identified so far from the biodiversity, it is obvious that new biological activities of known metabolites can be still put to light. Some of these known metabolites have only been structurally elucidated but never evaluated biologically. Among those that have already been engaged in one or even more specific bioassays, it is certain that a wide majority have not yet revealed all their biological potential, just because it remains impossible to test every compound on every known target in a reasonable amount of time. Furthermore, most *in vitro* bioassays are not relevant to *in vivo* or clinical conditions regarding either the activity or bioavailability of the compound under examination, not to mention that a metabolite with a specific activity in the pure state may show a different activity when present within a mixture

Innovative strategies are therefore needed to significantly reduce the timeline of bioactive natural product discovery.

All evidence suggests that the most promising workflows for the chemical profiling of natural products should involve, at least in part, a dereplication procedure. Historically, the first definition of the term "dereplication" was given by Beutler *et al.* in 1990 as "*a process of quickly identifying known chemotypes*" (Beutler et al. 1990). Their goal was to evaluate the activity of a range of terrestrial and marine plant extracts by using a simple phorbol dibutyrate (PDBu) receptor binding assay and to rapidly identify compounds responsible for this activity without investing time in traditional bioassay-guided fractionation or full structure elucidation procedures.

But what exactly lies behind the term "dereplication" more than twenty years later? Is this definition still relevant today within the context of natural product research? What does a dereplication process involve in terms of methods and techniques? For what purpose is such a strategy really useful? On which natural ressources is it currently applied? How much progress has been made in dereplication workflows over the last years?

In the present review, we would like to address these points by presenting a state-of-the-art report on the various dereplication approaches that have been used until now for the chemical profiling of natural products. A literature survey, covering the period 1990-2014, for the topic "dereplication" or "de-replication" in the Web of Science database revealed 358 hits. Upon closer examination, it emerged that the research areas covered by the matching studies not only include pharmacology/pharmacy, chemistry, or plant sciences, but also molecular biology, biotechnology, microbiology, and food science technology. Another important fact is the huge

4

increase of published items every year, particularly since 2012, as well as the exponential growth of annual citations (Fig. 1).



**Fig. 1.** Citation report obtained with the topic "dereplication" or "de-replication" from Thomson Reuters Web of Science. The latest 20 years up to December 2014 are displayed. (c) Thomson Reuters. All rights reserved

The literature survey was rigorously scrutinized in order to assess the function and usefulness of modern dereplication strategies for bioactive natural product discovery. Of course, the goal of this review is not to impose a definition or a specific way to perform natural product dereplication, but to highlight as objectively as possible that, depending on the natural resources under examination and on the final objectives, dereplication does not correspond to a single, clearly defined step-by-step procedure. In the 358 references found, 14 not dealing with natural products and 30 meeting abstracts were not taken into account. After a brief outline of the most common separation and analytical tools used for natural product dereplication, the critical role of natural metabolite databases and microbial strain libraries will be discussed. The studies covered by the literature survey have been organized in five distinct dereplication categories (annotated from DEREP1 to DEREP5), all involved in the search for novel bioactive substances of natural origin, but each following a particular workflow based on different starting materials, different separation and/or analytical procedures and different structural elucidation approaches.

A comparison chart summarizing these five categories (Table 1) shows that natural product dereplication can be performed as an untargeted workflow for the rapid identification of the major compounds whatever their chemical class in a single natural sample (DEREP1). Frequently, this first strategy is also directly combined with biological assays to accelerate bioactivity-guided fractionation procedures (DEREP2). In other cases dereplication is fully embedded in untargeted metabolomic studies for the chemical profiling of natural extract collections (DEREP3) or for the targeted identification of predetermined compounds (desired or undesired) in natural samples (DEREP4). Finally a distinct dereplication category mainly based on gene-sequence analyses is increasingly used for the taxonomic identification of microbial strains (DEREP5).

**Table 1.** Summary of the particular features distinguishing from DEREP1 to DEREP5

 dereplication workflows.

	DEREP 1	DEREP 2	DEREP 3	DEREP 4	DEREP 5
Goal	Identification of the major compounds in a single extract	Acceleration of activity- guided fractionation	Chemical profiling of crude extract collections	Chemical profiling of target compounds	Taxonomoic identification of microbial strains
Targeted chemical class?		No		Yes	No
Biological assays ?	Independent	Yes, systematically		Independent	
Samples	Single	extract	Extract collection	Single extract or collection	Extract collection
Fractionation step ?	In most cases Yes	Yes	No (direct sample analysis)	In most cases No	No (direct sample analysis)
Analytical tools	LC/M	IS, GC/MS and/or ]	NMR	Mostly LC/MS	Gene- sequencing

Computer tool and statistics for data treatments	May include	Yes, systematically	May include	Yes, systematically
Identification	Metabolite	database		Gene database

#### **Technical considerations**

As dereplication refers to the rapid identification of known secondary metabolites, it is obvious that reliable, robust, rapid and sensitive analytical methods are required to perform an efficient dereplication procedure. With the remarkable developments of separation sciences, spectroscopic techniques, and high-throughput analytical technologies, valuable resources now exist to structurally characterize natural products, not only as pure compounds, but also within complex mixtures such as crude plant extracts or microbial culture media. This review will start by presenting the most common analytical tools used for the dereplication of natural products. Since many detailed reviews are available for each technique, we will only briefly describe their strengths and weaknesses and focus on their benefits when applied to natural product dereplication.

#### Analytical chromatographic systems and hyphenation possibilities

High performance liquid chromatography (HPLC) remains by far the most commonly used technique to separate natural compounds at the analytical scale. HPLC is robust, easy-to-use, and applicable to a wide diversity of compounds, even starting from materials of complex chemical composition. More recently, ultra-high pressure liquid chromatography (UHPLC) has enabled a remarkable decrease of separation times while improving sensitivity, resolution, and reproducibility as compared to conventional HPLC. Both techniques, referred to as "LC" in the present paper, are currently applied to natural product analysis for various purposes including quality control, chemical characterization of crude plant extracts or high throughput

fingerprinting (Wolfender et al. 2010, Eugster et al. 2011). Thin layer chromatography (TLC or HPTLC) is another chromatographic technique frequently used for the preliminary analysis of natural extracts or for the rapid identification of known compounds within simplified mixtures obtained after the prefractionation of a crude natural sample (Hook et al. 1997, Hook 1998, Tan et al. 2011). TLC-based bioautographic methods also enable the rapid localization of biologically active metabolites directly on TLC plates (Hostettmann et al. 2005, Adhami et al. 2013) High speed counter current chromatography (SFC), centrifugal partition chromatography (CPC), supercritical fluid chromatography (SFC) or capillary electrophoresis (CE) are less commonly used analytical-scale separation system, and therefore will not be detailed here.

Among post-chromatographic hyphenation possibilities, simple ultraviolet (UV) or photodiode array (DAD) detectors remain limited to the interpretation of peak retention times or UV spectral fingerprints, but can nevertheless be employed in dereplication strategies for both identification and quantification purposes by using standard molecules as references (Wolfender 2009). With the help of chemometric tools, LC/UV-DAD data can be processed by chromatographic band deconvolution, resulting in a significant improvement of chromatographic resolution. This method was recently used to identify several Cglycosylflavone isomers in the traditional herbal medicine *Jatropha gossypifolia* (Pilon et al. 2013). However, due to the very limited structural information recovered from simple UV or DAD detection systems, other more powerful instruments are generally required to assess the structural diversity of natural products. Among them, mass spectrometry (MS) and nuclear magnetic resonance (NMR) described in the next section remain by far the favourites.

#### MS and NMR-based detection in natural product dereplication workflows

Mass spectrometry is a sensitive, rapid and accurate high throughput detection technique frequently used for the dereplication of natural products. The characterization of natural metabolites is established on the basis of exact mass, elemental composition, adducts and fragmentation patterns. Due to their high sensitivity, MS-based methods also enable the rapid detection of trace-level compounds, meaning that only small amounts of materials is required for successful analyses. LC coupled to high-resolution mass spectrometers such as Time-of-Flight (TOF), Fourier Transform (FT) or Orbitrap devices currently constitute the most powerful "high-throughpout screening" platforms for the on-line identification of metabolites in natural resources (Strege 1999, Wolfender et al. 2000, Shin and van Breemen 2001, Harrigan and Goetz 2005, Bitzer et al. 2007, Moss et al. 2007, Sashidhara and Rosaiah 2007, Nielsen, Mansson et al. 2011, Sarker 2012, Smyth et al. 2012, Carter 2014). Regarding hydrophobic or volatile small molecules, GC/MS is a more appropriate coupling, as demonstrated for instance for the dereplication of fatty acids (Stavri et al. 2004) or flavor and aroma constituents (Molyneux and Schieberle 2007).

Although MS is recognized as a key technology for the identification of natural products, some pitfalls still exist and these problems have to be solved to make MS-based dereplication methods more efficient (Cech and Yu 2013). The major drawback arises from the important variabilities in the raw datasets obtained from one mass analyzer to another, which strongly hampers the creation of exchangeable MS/MS databases. The ionization processes taking place at the input of MS systems to desolvate and charge the analytes are also particularly critical because they vary with the type of ion source (electrospray or atmospheric pressure chemical ionization, for example), and may also vary through ionization suppression or enhancement under matrix effects. A high heterogeneity has been reported after the construction of a Collision-Induced-Dissociation MS/MS spectral library of ubiquitous flavonoids using either hybrid quadrupole time of flight (Q-TOF) or Ion Trap (IT) under various CID energy conditions

(Wolfender et al. 2000). Perfect control of the parameters used to generate informative MS/MS spectra is thus needed for an efficient use of MS detection in the dereplication of natural products.

The interpretation of MS data is another critical issue that limits the speed of dereplication strategies. Efforts are currently underway to develop the computerized treatment of mass spectral data. Software dedicated to the pre-processing of LC-MS data are now available and very useful for peak picking, ion extraction, organization and classification of data. They include for instance Bruker1 Data Analysis and Bruker1 Profile Analysis for Bruker, MassHunter for Agilent, MarkerLynx for Waters or publicly available software such as XCMS, MZmine, and METIDEA which can handle data from different instruments. Among the computer tools that assist metabolite identification, four data mining software are for instance described in a recent review (Hufsky 2014): MetFusion which uses substructures to assess spectral and chemical similarities, ISIS which is based on the prediction of fragmentation patterns for spectra comparison, FingerID which compares molecular structures after prediction of structural features, and FT-BLAST which is based on the use of a fragmentation tree database.

Molecular networking is also a promising computer-based approach to visualize and organize tandem mass spectrometry datasets and to automate database search for metabolite identification within complex mixtures (Garg et al. 2015). Such a tool relies on the observation that structurally similar metabolites share similar MS/MS fragmentation patterns. After collection of MS/MS spectra from a natural sample, the method consists in the construction of a molecular network by measuring the correlations within the produced dataset and establishing a Cytoscape visual representation of the chemical similarities between metabolites. Because structurally similar metabolites share similar fragmentation patterns, molecular families tend to cluster together within a network (Nguyen, Wu et al. 2013, Yang, Sanchez et al. 2013). The

chemical investigation of natural resources by molecular networking has already proven to be a very useful complement to current MS-based dereplication strategies, for instance to highlight the chemical diversity among marine cyanobacteria (Winnikoff, Glukhov et al. 2014), to automatically detect structurally related nonribosomal peptides in filamentous fungi (Klitgaard, Nielsen et al. 2015), or to investigate interkingdom molecular interactions and metabolic exchange processes within microbial communities (Moree, Phelan et al. 2012).

Another promising data mining workflow based on MS<sup>2</sup> precursor lists and targeting only signals related to bacterial metabolism was recently developed to characterize myxobacterial secondary metabolites (Hoffmann et al. 2014). Even direct MS infusion can be used for natural product dereplication if assisted by computer tools. Such an approach was applied to the chemotyping of filamentous fungi strains from culture collections, and to subsequently classify unknown samples, thus avoiding redundancies in the selection of samples to be further investigated (Larsen 2005).

Nuclear Magnetic Resonance (NMR) is the other predominant detection technique used for the dereplication of natural products. It remains by far the most efficient to unambiguously elucidate complex structures of individual small molecules. With the advent of high field magnets, capillary and cryogenic probes, the lower sensitivity of NMR as compared to MS-based analytical methods has been progressively counterbalanced, and even the structure elucidation of minor compounds within mixtures becomes possible (Hu et al. 2008). In addition, sophisticated hardware and software products have been introduced in recent years to promote high-throughput NMR analyses, such as NMRbot Python scripts (Clos et al. 2013). As described in a recent review, these technical improvements along with hyphenation possibilities and their combination with computational treatments is progressively bringing NMR to the front of the powerful resources used in natural product dereplication strategies (Halabalaki et al. 2014).

It should nevertheless be emphasized that solvent and pH effects induce significant chemical shift variations across samples and are important drawbacks of NMR-based methods. These effects can strongly influence the efficiency of natural product dereplication procedures, as demonstrated for instance in a study investigating the molecular structures of caffeoyl quinic acid derivatives (Pauli et al. 1999). The limited spectral dispersion and the complexity of signal patterns, particularly in <sup>1</sup>H NMR spectra are also a major NMR challenge. In this sense it was pointed out in a recent paper that unambiguous <sup>1</sup>H NMR analyses for dereplication purposes require adequate precision when reporting chemical shifts or <sup>1</sup>H-<sup>1</sup>H coupling constants from <sup>1</sup>H spectra, what is not so that easy when working with spectra of natural product mixtures (Pauli 2014). Diffusion ordered spectroscopy (DOSY) was proposed as an efficient alternative to simplify interpretation of NMR spectra of natural mixtures (Williamson et al. 2000, Stessman et al. 2002). It has been shown for instance that 1D-DOSY when combined with 1D or 2D NMR experiments enables the acquisition of full spectral data of high molecular weight polysaccharides directly from crude hot water extracts of mushroom species without the need of any purification step (Politi et al. 2006) Although natural organic compounds also contain carbon atoms, the use of <sup>13</sup>C NMR for the analysis of natural extracts remains largely underutilised, mainly because the low abundance of the <sup>13</sup>C isotope (1%) and its low gyromagnetic ratio (25 % of that of <sup>1</sup>H) considerably reduce its detection sensitivity. Yet <sup>13</sup>C NMR presents several strong advantages for the analysis of natural mixtures: carbon atoms constitute a significant part of all organic molecules, and each <sup>13</sup>C position in a molecular structure corresponds to a single resonance on a broadband <sup>1</sup>H decoupled <sup>13</sup>C NMR spectrum. In addition, the <sup>13</sup>C NMR spectral width is significantly higher than that of <sup>1</sup>H (220 ppm for <sup>13</sup>C and 12 ppm for <sup>1</sup>H), what also significantly reduces the occurrence of signal overlaps. Here again with the technical improvements of NMR spectrometers and the emergence of promising methodologies such as dynamic nuclear polarisation, it becomes possible to acquire <sup>13</sup>C spectra of metabolite mixtures with high resolution, good sensitivity, and in a reasonable time. In fact, an increasing amount of work using <sup>13</sup>C NMR for the analysis of natural products is emerging (Clendinen et al. 2014, Hubert et al. 2014, Yona et al. 2015).

Particular attention must also be paid to LC/NMR or LC/SPE/NMR methods, which combine the separation performance of liquid chromatography, the concentration effect of SPE, and the structural information provided by NMR. In the case of LC/SPE/NMR, the main advantage is that peak trapping on SPE cartridges significantly increases analyte concentration and thus provides access to 2D NMR data in a short time. The use of LC/NMR has rapidly expanded over the last years for the dereplication of natural products, especially for the direct comparison of metabolite profiles in small extracts using <sup>1</sup>H NMR before further investing in more laborious preparative-scale isolation processes. Such strategy greatly helps to avoid re-isolation of already known extract constituents. Several reviews published since the mid 1990s have detailed LC/NMR technical developments and their applications in the field of natural product research (Bobzin et al. 2000, Bobzin et al. 2000, Wolfender et al. 2001, Wolfender et al. 2003, Jaroszewski 2005, Jaroszewski 2005, Wolfender et al. 2013).

#### The crucial role of natural product databases and computer tools

The success of a dereplication procedure to achieve the rapid characterization of known compounds greatly depends on the availability and quality of natural product databases or strain libraries. Many commercial or public databases of low molecular weight metabolites have been developed over the last 20 years (Corley and Durley 1994, Lopez-Perez et al. 2007, Blunt and Munro 2013, Valli et al. 2013) The more comprehensive databases include the *Dictionary of Natural Products* (DNP) containing over 260,000 molecular structures, the *Dictionary of Marine Natural Products* containing around 48,000 molecular structures drawn from marine

organisms, the *ACD/Labs NMR* and *ACD/Spectrus* databases providing structural and spectral data of many thousands of synthetic and natural compounds, the *UCSD Natural Products Database* containing a large repository of microorganisms and exosymbionts, the *Human Metabolome Database*, *Napralert*, the *NuBBE* database dedicated to the Brasilian biodiversity, the UniProtKB/Swiss-Prot protein sequence database, *KnapSack* which is a comprehensive plant species-metabolite relationship database, and *MarinLit*, *AntiMarin*, *AntiBase* or *Seaweed Metabolite* databases dedicated to marine and/or microbial molecular structures. This is not to mention all the locally-built libraries fragmented and scattered in many laboratories worldwide. There is a crucial need today to expand and standardize these natural products databases.

A number of studies found in the literature survey were performed with the sole objective of database construction and implementation. Many of them were based on molecular structures, while others were focused on MS fragmentation patterns, bioactivity or taxonomic data (Fredenhagen et al. 2005, Konishi et al. 2007, Liu et al. 2008, Overy et al. 2008, Rojas-Cherto et al. 2012).

In addition to databases, computer tools and multivariate statistics have considerably improved the automation of analytical data processing and have substantially contributed to the current enhancement of natural product dereplication strategies (Hansen et al. 2005, Wolf and Siems 2007, Jarussophon et al. 2009, Ng, Bandeira et al. 2009, Khosrokhavar et al. 2010, Wolfender et al. 2010, Chlipala et al. 2011, Eugster et al. 2014, Kokkotou et al. 2014). Strong advances in the area of Computer-Assisted Structure Elucidation (CASE) systems, which combine artificial intelligence software with spectroscopic data to automatically generate molecular structures, have also dramatically accelerated structure elucidation procedures and have improved the reliability of dereplication results (Jaspars 1999, Steinbeck 2004, Elyashberg et al. 2009). Finally, spectral pattern recognition strategies based on the organization of spectral data according to chemical similarities are also powerful complements to natural product dereplication workflows. In this case spectroscopic data of metabolite mixtures are required along with spectra obtained from known standards, well-characterized organisms or prediction tools, preferably organized into databases. The utility of pattern recognition tools for natural product dereplication was for instance demonstrated for the chemical characterization of a diverse array of marine and terrestrial microbial samples using a MS-based method (Yang et al. 2013), or for the chemical characterization of plant extracts using <sup>13</sup>C NMR combined to hierarchical clustering analysis (Hubert et al. 2014).

#### Dereplication and metabolomics: Where to draw the line?

Before entering the literature survey, some thought should be given to the term "chemical profiling" as well as to the obvious connection between dereplication and "metabolomics". Of course each of these two terms alone deserves detailed discussion, as their use can be ambigous depending on the context. For the sake of clarity, we would like to define the meaning of "chemical profiling" within the context of the present review, and also to highlight that a range of metabolomic studies are fully associated with natural metabolite dereplication workflows.

The term "chemical profiling" will be defined here in a very broad sense as the systematic detection and/or identification and/or quantification of a range of natural compounds. Depending on the final purpose and on the analytical technique available, chemical profiling can be performed as a fingerprint analysis or as a targeted profile. A fingerprint analysis represents a global analysis where all detected metabolites are not necessarily identified. In a targeted chemical profiling approach, a predefined number of compounds or a particular chemical class of compounds is investigated and the molecular structures are identified.

Metabolomics, which is defined as the qualitative and quantitative analysis of the whole set of low molecular weight metabolites present in a biological system (Fiehn 2002, Zhang et al. 2012), also falls within the scope of a chemical profiling. However, more specifically, metabolomics is an interdisciplinary field that combines high resolution analytical systems, multivariate statistics and data mining tools, chemical and biological knowledge, and sometimes metabolic network modeling in an attempt to understand metabolic pathways, genefunction relationships, or states of an organism in response to environmental changes, drug perturbations, phylogeny, genotypic or phenotypic variabilities. Metabolomics workflows are strongly context-dependent and have been used over the last years in so many different fields that a range of experimental design methodologies have been developed with specific lines regarding sample preparation, data analysis or metabolite identification strategies. To keep it simple, we will only distinguish "targeted" from "untargeted" metabolomics studies. Typically, a targeted metabolomic approach is used to analyze a set of defined metabolites in a sample, while a much more comprehensive "untargeted" metabolomics approach consists of the detection and/or quantification of as many metabolites as possible without a priori knowledge of metabolite targets. In this latter case, samples and metabolites are generally classified according to spectral patterns. In the field of natural product research, both targeted and untargeted metabolomic approaches have already demonstrated strong potentialities, mainly for the quality control of herbal medicines, for the determination of chemical markers in plant species, or to assess the correlations between bioactivity and composition (Fiehn 2002, Yuliana et al. 2011, Sheridan et al. 2012, Zhang et al. 2012, Cox et al. 2014, Seiber et al. 2014). Of course the use of metabolite databases for the identification of the metabolites was a key aspect of all of these studies, and although not commented on explicitly, a dereplication process was in fact involved in the metabolomics study every time that a known metabolite was identified with the help of a database. It must be mentioned here that in dereplication, metabolite identification is often only a putative identification. Several articles have suggested ways to evaluate the level of confidence in metabolite identification (Dunn, Erban et al. 2013). For instance the Chemical Analysis Working Group of the Metabolomics Standards Initiative (http://msi-workgroups.sourceforge.net) has defined four different confidence levels, with level 1 corresponding to confidently identified compounds by comparing multiple physicochemical properties of a pure standard to those of the metabolite of interest, level 2 corresponding to putatively annotated compounds, level 3 corresponding to putatively annotated compound classes and level 4 corresponding to unknown compounds (Sumner, Amberg et al. 2007, Creek, Dunn et al. 2014). Implementation of such levels, mainly in the field of metabolomics, would also be useful for the natural product research community.

#### Dereplication of natural products: one concept involved in different strategies

Rapid identification of the major compounds in a single natural extract regardless of chemical class (DEREP1)

A very common way to perform dereplication of natural products today is to establish the chemical profile of a single natural resource using a hyphenated analytical system and to directly compare the obtained spectroscopic data with libraries of natural metabolites. This dereplication approach will be annotated "DEREP1" in the present review and is illustrated in Fig. 2.



**Fig. 2.** DEREP1: an untargeted dereplication workflow aiming at the rapid identification of the major known compounds whatever their chemical class in a single natural sample

The starting natural resource is most often fractionated before the analytical step and the dereplication procedure can be achieved independently from bioassay experiments. In this way, a global view of the main chemical classes present in a natural species can be easily obtained and theoretically the major known compounds rapidly identified. Among the published works mentioning the term "dereplication" between 1990 and 2014, 50 studies summarized in Table 2 were based on a DEREP1 workflow (16 % of the total examined articles). More than two-thirds of these studies were carried out on plant extracts, while a minority delt with marine sponges, brown algae and cyanobacteria extracts. This distribution is logical considering that databases containing spectroscopic data of plant-derived metabolites are substantially more developed than those related to less explored marine or microbial species. The analytical tools involved in these studies were usually performed by directly entering spectral data (exact mass or molecular formula in MS, chemical shifts in NMR, UV spectra or log P-based estimation of the chromatographic retention time in LC/DAD) into a database for automatic search. In several studies the strength of the procedure was reinforced by combining the analytical step with

computational treatments in order to detect spectral patterns before searching molecular structures into metabolite databases (Boudreau et al. 2012, Hubert et al. 2014).

It can also be noted from Table 2 that, even if flavonoids and phenolic acids remain by far the most investigated compounds, a diversity of other chemical classes including anthrones, iridoids, terpenoids or fatty acids have also been successfully identified in these DEREP1 studies. Another advantage is that although dereplication does not provide original information on novel chemical structures, previously known compounds can still be identified in genus or species in which they are not *a priori* expected. For instance, flavonoid glycosides already known in a range of plant species were identified for the first time in the genus *Lasiopetolum* by using a LC/NMR-based DEREP1 procedure (Timmers and Urban 2011). Of course, the efficiency of such dereplication approach strongly depends on the quality of spectral libraries. As mentioned above, significant efforts are currently being to develop and especially to standardize comprehensive databases containing structures and spectroscopic data (MS or NMR) of natural products.

Identified chemical classes and natural sources	Analytical technique	Reference
Representative set of 46 plant and microbe-derived	ESI-MS, API-MS	Zhou and
metabolites		Hamburger 1996
An alkaloid and a sesquiterpene previously purified	NMR, computer-	Bradshaw et al.
	assisted identification	2001
	using the DNP	
Three pentaketides and four fungal pigments from a deep	1D and 2D NMR	Gautschi et al.
water marine-derived fungal culture		2004
Ent-labdane diterpene glycosides from polar extracts of	LC/MS, LC/MS <sup>2</sup> ,	Waridel et al.
Potamogeton species	LC/UV, LC/NMR	2004
Enantiomeric 1-acetoxychavicol acetates and carvones	Chiral Lanthanide	Jaki et al. 2004
	shift reagents in NMR	
Flavonol glycosides and cardenolides from flower, leaf, root,	LC/DAD/SPE/NMR	Clarkson et al.
and stem extracts of Kanahia laniflora		2005

**Table 2.** Dereplication strategies published from 1990 to 2014 using a DEREP1 workflow(from oldest to most recent).

Aspirochlorine and two new antifungal derivatives from	LC/UV/ELSD/MS,	Klausmeyer et al.
Aspergillus flavus	NMR	2005
Cyclic peptides from a crude fungal extract	LC/UV, HSCCC	Dalsgaard et al. 2005
Senkyunolide A, butylphthalide, neocnidilide, Z-ligustilide	LC/UV, LC/MS,	Zschocke et al.
and several phthalide dimers from the TCM Ligusticum	LC/NMR	2005
chuanxiong		
Ursene triterpenes from Diospyros dendo with antibacterial	Capillary-scale NMR	Hu et al. 2006
activity against Pseudomonas aeruginosa		
Panel of 179 natural standard molecules	<sup>1</sup> H and <sup>13</sup> C NMR	Dunkel and Wu
		2007
Huperzine A and a Lycopodium anti-Alzheimer drug lead	<sup>1</sup> H NMR, spectral	Niemitz et al.
alkaloid	prediction	2007
Naphtodianthrones, phloroglucinols, flavonoids and	LC/DAD/SPE/NMR,	Tatsis et al. 2007
phenolic acids from Greek Hypericum perforatum	LC/UV/ESI-MS	
Several bioactive compound from the Chinese herbal	LC/DAD/MS,	Liu et al. 2008
formula Qi-Xue-Bing-Zhi	computer-assisted	
	data processing	
	(WiseProcessor) and	
	library	
Flavonoids, sesquiterpene lactones and phenolic acids from	LC/DAD/MS and	Gobbo-Neto and
Brasilian Lychnophora ericoides leaves	$MS^2$	Lopes 2008
Pentaprenylated <i>p</i> -quinol from an antitumour extract of the	LC/MS, LC/NMR	Dias and Urban
marine sponge Dactylospongia sp.		2009
Bastadins, 19-Hydroxyaraplysillin-I-N <sup>20</sup> -Sulfamate and	FTICR/MS, MS <sup>n</sup>	Motti et al. 2009
Araplysillin-I-N <sup>20</sup> -sulfamate from the marine sponge		
Lanthella flabelliformis		
Oxy-polyhalogenated diphenyl ethers from <i>Dysidea</i> sponges	NMR, X-ray crystal	Calcul al. 2009
Potent antiplasmodial quinolinone alkaloids from Haplophyllum acutifolium	LC/DAD/MS/SPE/N MR	Staerk et al. 2009
Fatty acids from the antifouling marine sponge Cholla	TLC, GC/MS, NMR	Castellanos et al.
delitrix		2010
Chlorinated cylindrocyclophanes from a Nostoc	LC/UV/MS	Chlipala et al.
cyanobacteria collected in a parkway of Chicago with		2010
antiproliferative activity		
Five cyclic tetrapyrrolic photosensitisers from <i>Phaeanthus</i>	TLC, LC	Tan et al. 2011
ophtalmicus leaves		
Spiro compounds and tracheloside from Carthamus	HPLC/DAD/MS/SPE/	Johansen et al.
oxyacantha	NMR	2011
719 microbial natural product and mycotoxin standards	LC/DAD/TOF-MS,	Nielsen et al. 2011
	MS/MS under	
	different ion-source	
	settings	
Linear polyketides and tubercidin from Actinopolyspora	1D and 2D NMR	Zhao et al. 2011
erythraea		
Quantification of structurally diverse standard natural	ELSD	Adnani et al. 2012
products		

Tiliroside derivatives (on-line) and flavanoid glycosides	On-line LC/NMR,	Timmers and
(off-line) from Lasiopetolum macrophyllum	off-line LC, NMR,	Urban 2011,
	MS	Timmers and
		Urban 2012
Alkaloids, amides or esters of hydroxycinnamic acid and	LC/MS	Nikolic et al. 2012
betain from Cimicifuga racemosa (black cohosh)		
Phenolic allopyranosides, amide allopyranosides, and	LC/NMR, LC/MS	Yim et al. 2012
phenolic compounds from the rhizomes of Cimicifuga		
heracleifolia		
Stem-specific metabolites including three stilbenoids from	HPTLC/MS	Cakova et al. 2012
Vanda coerulea (Orchidaceae)		
Beilschmiedic acid derivatives from the leaves of a	Micro-CryoProbe	Williams et al.
Gabonese Beilschmiedia species	NMR	2012
Hectochlorin derivatives and several jamacamides from the	$MS^2$	Boudreau et al.
cyanobacteria Moorea producens		2012
Iridoid and triterpenoid glycosides from Premna fulva leaves	LC/DAD/MS	Niu et al. 2013
and stems		
Phenolic glycosides, dimeric phenylpropanoid glucoside,	LC/DAD/MS, NMR	Abbet et al. 2013
saponins, and fatty acids from Phyteuma orbiculare leaves		
Two known naphthocoumarins and one new	HPLC-PDA/LC-MS	Jain et al. 2013
naphthocoumarin from Streptomyces sporoverrucosus	combined to DNP, 2D	
	NMR	
Resorcinol and two polyene derivatives from the brown	LC/NMR	Urban and
algae Cystophora tondosa		Timmers 2013
Spiro compounds from Carthamus oxyacantha (wild	LC/NMR	Johansen et al.
safflower) and griseofulvin and analogues from the		2013
endophytic fungus Penicillum namyslowski		
Leporizines from an Aspergillus sp. strain	NMR	Reategui et al.
		2013
Phenolic glycosides, monoterpene lactone, spermine	LC/DAD/MS,	Abbet et al. 2014
derivatives and fatty acids from <i>Cirsium spinosissimum</i>	microprobe NMR	
Isomeric butanoates and pentanoates of long-chain 1-	GC/MS	Radulovic et al.
alkanols from the essential oil of <i>Scandix pecten-veneris</i>	12 C N 1 C A	2014
Triterpenes, saponins, phenolic acids, flavonoids and ellagic	<sup>13</sup> C NMR, HCA	Hubert et al. 2014
acid derivatives from the bark of <i>Anogeissus leiocarpus</i>		YZ'1 1 1 1 1
Polyketides, non-ribosomal peptides, terpenes,	LC/DAD/QIOF-MS,	Kildgaard et al.
meroterpenoids and four novel isomers of the known	MS/MS	2014
anticancer compound asperphenamate, from marine-derived		
strains of Aspergillus, Peniculium, and Emericellopsis	LOME	View language of all
Hydroxylated fatty acids, antimycin compounds and three	LC/MS	Viegelmann et al.
butenolides from a sponge-derived <i>Streptomyces</i> sp. with		2014
Indole and norththe suine and derivatives from a north	MC NMD	Jansan at al. 2014
muone and naphilioquinone derivatives from a novel		Jansen et al. 2014
Diplosporin corilovin melloin and coistating derivatives		Ibrohim at al
from a <i>Yularia</i> sp. isolated as an and apply to from a surface		2014
storilized Concord graph leaf (Vitig Jahrunger)		2014
stermzed Concord grape lear ( <i>vitis labrusca</i> )		

Dioxomorpholine, okaramine, and aflavinine derivatives and	LC/DAD/MS, NMR	Petersen et al.
three novel structures of mixed biosynthetic origin named		2014
aculenes A-C from the black filamentous fungi Aspergillus		
aculeatus		
Four macrotetrolides homologous to nonactin and three	MS, LC/MS <sup>2</sup>	Crevelin et al.
related linear dimers from Streptomyces sp.		2014
Six major depsides from the lichen Pseudevernia furfuracea	<sup>13</sup> C NMR, HCA	Oettl et al. 2014
Ten known compounds, including angucycline,	NMR	Dashti et al. 2014
diketopiperazine and $\beta$ -carboline and three novel derivatives		
produced in the culture medium of Actinokineospora sp.		
EG49 grown in co-culture with Nocardiopsis sp.		

Dereplication workflows involved in bioactivity-guided fractionation procedures (DEREP2)

A widely used strategy in the search for new bioactive molecules among natural products consists of the so-called bioactivity-guided fractionation procedures, in which pharmacological or biological assays are performed to target the isolation of active constituents. They were initially developed to avoid wasting time and resources in considering inactive, thus arbitrarily implying "uninteresting" compounds, and focusing only on the fractions or metabolites with a predefined biological activity (Agarwal 2014). Once fractions obtained from a crude extract have been revealed active by biological assays, classical separation and analytical techniques are applied to isolate and identify the individual active substances (Duarte et al. 2012, Smyth et al. 2012). Such procedures are currently very common in natural product research, and a range of chemical, enzymatic, and *in vitro* biological tests (antioxidant, antimicrobial, antifungal, anti-inflammatory...) are routinely available in many laboratories (Hostettmann et al. 2001, Cos 2006).

However, the identification of new compounds by bioactivity-guided fractionation procedures requires a multi-step workflow, and often a great deal of work is done to purify the bioactive fractions and finally discover previously known compounds (Agarwal et al. 2014).

Consequently, dereplication also constitutes a key approach to overcome this frequent issue of known compound rediscovery in bioactivity-guided fractionation procedures.

In the literature survey, 52 studies (17 % of the total examined articles) were dedicated to the identification of natural metabolites on the basis of bioactivity-guided fractionation procedures. These approaches will be annotated DEREP2 in the present paper. It is obvious that DEREP1 and DEREP2 workflows are not that different, except that DEREP2 ones are systematically focused on active natural samples as determined by biological assays. These studies are summarized in Table 3 and their common global workflow is illustrated in Fig. 3.



**Fig. 3.** DEREP2: A dereplication workflow for the acceleration of bioactivity-guided fractionation procedures

Among these studies, a vast majority was again focused on plant extracts and a minority on marine and microbial extracts. In some cases, dereplication was performed at an early stage in an attempt to rapidly distinguish known constituents from those presenting novel or unusual spectroscopic features before undergoing multi-step isolation procedures. From an analytical point of view, DEREP2 workflows are most often based on LC/MS<sup>n</sup> and/or NMR to analyze biologically active fractions obtained after the pre-fractionation of the crude extract. As

described in Table 3, the resulting molecular masses, fragmentation patterns or NMR chemical shifts are then compared to data from the literature or submitted to commercially available or in-house natural product databases for the identification of known compounds. Analytical data that does not correlate with any known compound thus theoretically corresponds to novel molecules that can be further investigated and ultimately find application as new active substance. For example, this strategy was thoroughly described in a study investigating antibacterial substances in the leaves of *Melicope vitiflora* by using an *in vitro* biological assay in combination with a MS-based dereplication method (O'Donnell et al. 2009). More indirectly, dereplication was also reported useful when combined with bioactivity-guided fractionation procedures to rapidly set aside undesired constituents. For instance, the dereplication of sulphated polysaccharides was achieved in two studies investigating new potent anti-HIV compounds, because many of them were already extensively described in the literature as having an anti-HIV activity and thus were not considered as interesting compounds (Beutler et al. 1993, Harnett et al. 2005). In this way, the tedious isolation of undesired constituents could be avoided, so that isolation, structure determination, and pharmacological investigations could be carried out only on bioactive and novel compounds.

It must be pointed out that most bioactivity-guided fractionation procedures are based on the theory that the activity shown by a mixture of metabolites results directly from the sum of activities of individual metabolites. However, it is becoming evident that the biological effects of a natural extract most likely result from a multi-component synergism (Pavana et al. 2009, Wagner 2011, Yuliana et al. 2013). Another limitation reported for bioactivity-guided fractionation is the irrelevance of most *in vitro* bioassays as compared to *in vivo* or clinical conditions regarding the efficacy and bioavailability of the samples under examination (Houghton 2007). Potentially interesting compounds may also be missed just because they are not active on the selected biological assays. Research strategies for the discovery of novel

bioactive molecules among natural products are thus progressively re-evaluated to consider more than one active constituent within natural resources (Wang et al. 2012).

Table 3	. Dereplication	strategies	published	from	1990 to	o 2014	using a	DEREP2	workflow
(from old	lest to most rece	ent)							

Natural materials	Biological assays	Compounds	Analytical	Ref
		identified	technique(	
			<b>s</b> )	
Lyngbya majuscula	Phorbol dibutyrate (PDBu)	Debromoaplysiatoxin	HPLC/UV	Beutler et al.
(filamentous	receptor binding assay	in Lyngbya majuscula		1990
cyanobacteria) and		and a complex of		
Croton cuneatus		potent phorbol esters		
(plant)		in Croton cuneatus		
Marine invertebrates	HIV inhibition test	Sulphated	Precipitatio	Beutler et al.
		polysaccharides (for	n method	1993
		removal)		
Thevetia ahouia	Cytotoxicity against a panel of	Cardenolide	UV, IR,	Decosterd et
(wood)	human cancer cell lines	glycosides	MS, NMR	al. 1994
Marine sponge Agelas	In vitro radioreceptor binding	Agelasines	NMR	deVries et al.
axifera	assay on protein kinase C			1997
Azadirachta excels	Cytotoxicity against KB cell	Meliacin-type	LC/ESI-	Cui et al.
(stem bark)	lines	limonoids	MS	1998
Semecarpus	Cytotoxicity against a panel of	Alkenyl catechols	HPLC/ESI-	Shin et al.
anacardium	human cancer cell lines		MS	1999
Five crude plant	Radioligand Receptor Binding	Galanthamine,	CPC, TLC	Ingkaninan et
extracts	Assays; AChE Inhibitory	theobromine, caffeine,		al. 2000
	Activity; in vitro anti-	fatty acids, phenolic		
	asthmatic assay	acids and flavonoids		
Acnistus arborescens	Cytotoxicity against a panel of	Withanolides	NMR, MS	Minguzzi et
(leaves)	human cancer cell lines			al. 2002
Kielmeyera	Cytotoxicity against KB cell	Coumarins	LC/MS	Scio et al.
albopunctata (bark)	lines			2003
Plant extract spiked	Continuous-flow enzymatic	Phosphodiesterase	LC/fluoresc	Schenk et al.
with 2 naturally	assay	(PDE) inhibitors	ence	2003
occurring PDE				
inhibitors				
Artocarpus kemando	Cytotoxicity against KB cell	Prenylated flavonoids	HPLC/ESI-	Seo et al.
(stem bark)	lines and DNA strand-scission		MS	2003
	activity			
Plant extract spiked	Continuous-flow enzymatic	Phosphatase inhibitors	LC/MS	Schenk et al.
with tetramisole	assay		(on-line)	2003
In vitro marine culture	Antifungal activity against	Griseofulvin	ESI-IT/MS	Petit et al.
of the strain	Candida albicans		and MSn	2004

Penicillium waksmanii				
Zaleski				
Humulus lupulus	Antimycobacterial activity	Fatty acids	NMR,	Stavri et al.
-	against Mycobacterium		GC/MS	2004
	fortuitum			
Macrococculus	Cyclooxygenase-2 inhibition	Dibenzylbutyrolactone	NMR, MS	Su et al.
<i>pomiferus</i> (stems)	test	lignans and seven		2004
		known compounds		
Punica granatum	Estrogen receptor binding	Luteolin, quercetin and	LC/MS	van Elswijk
(peel)	assay	kaempferol	(on-line)	et al. 2004
Sutherlandia	HIV inhibition test	Sulphated	Precipitatio	Harnett et al.
frutescens and		polysaccharides (for	n method	2005
Lobostemon trigonus		removal)		
Antibacterial lead	Live/Dead bacterial viability	Nine known	Fluorescen	Singh 2006
compounds	kit containing two fluorescent	antibiotics and 14	ce	e
1	nucleic acid stains	novel lead compounds	microscopy	
Cleistopholis patens	Antibacterial activity against	Three new and five	capillary-	Hu et al.
1 1	Staphylococcus aureus	known acetylated	scale NMR,	2006
	1 2	oligorhamnosides	ESI-MS	
Six <i>Terminalia</i> species	<i>In vitro</i> antifungal activity ( <i>C</i> .	Non-polar antifungal	bioautograp	Masoko and
1	albicans, C. neoformans, A.	compounds	hy	Eloff 2005
	fumigatus, M. canis and S.	1	5	
	schenkii)			
Dracaena angustifolia	Anti-mycobacterial test	Three compounds	TLC	Case et al.
(leaves)	-	(ergosterol-5,8-		2007
		endoperoxide, linoleic		
		acid		
		and E-phytol)		
Alangium longiflorum	Inhibition of transcriptional	Tubulosine and two	NMR	Klausmeyer
	activation pathway in human	derivatives		et al. 2008
	cancer cell lines			
Ophiorrhiza	Inhibition of transcriptional	Camptothecin and	LC	Klausmeyer
trichocarpon	activation pathway in human	three analogues		et al. 2007
-	cancer cell lines	C C		
Parthenium hispitum	Inhibitory activity against	Five new and four	LC,	Hu et al.
	hepatitis C virus	known oxygenated	capillary-	2007
	-	hydroxy-	scale NMR	
		pseudoguaienolides		
Petiveria alliacea	Cytotoxicity against several	13 possible known	LC/MALD	Uruena et al.
	tumor cell lines	compounds	I-TOF-MS	2008
Distephanus	In vitro antiplasmodial assay	Chlorogenic acid	HPLC/DA	Pedersen et
angulifolius		analogs, steroid	D/MS,	al. 2009
		saponins and	SPE/NMR	
		sesquiterpene lactones		
Albany Molecular	In vitro activity against multi-	Mutactimycin E	NMR	Hopp et al.
Research Inc. (AMRI)	drug resistant Staphylococcus	-		2008
natural product library	aureus and counter screening			

	for cytotoxicity against the human HepG2 cell line			
Melicope vitiflora	Antibacterial activity against methicillin-resistant- <i>Staphylococcus aureus</i> and <i>Micrococcus luteus</i>	12 known compounds	TLC/ESI- MS <sup>n</sup>	O'Donnell et al. 2009
Cimicifuga racemosa	Serotonergic activity using a 5-HT(7) bioassay	Cimicifugic acids	NMR (structure- based spin- pattern analysis)	Goedecke et al. 2009
Myxobacterial extracts	Bacterial bioassay based on a whole-cell bioluminescent reporter gene assay	Inthomycin A	TLC, LC/MS, LC/NMR	Kreiss et al. 2010
Aegle marmelos (Bael tree)	Inhibition of transcriptional activation pathway in a human breast tumor cell lines	Two protolimonoids	NMR	Li et al. 2011
<i>Cacospongia</i> <i>mycofijiensis</i> (marine sponge)	Cytoskeletal profiling, cytotoxicity and antiparasitic activity	Twelve known and four new compounds	LC/MS/UV /ELSD	Johnson et al. 2011
<i>Syzygium polyanthum</i> leaves	Photocytotoxicity	Two new phloroglucinol derivatives and five known pheophorbides	Data not found	Har et al. 2012
Medicinal plants	<i>In vivo</i> zebrafish bioassays applied on microfractions	Anticonvulsant and antiangiogenic compounds	Microflow NMR, UHPLC/M S	Challal et al. 2012
<i>Diospyros peregrina</i> fruits	1,1-diphenyl-2-picryl-hydrazyl (DPPH) radical scavenging assay	Luteoline-4'-methyl- ether-7- <i>O</i> -glucoside and quercetin-3- <i>O</i> - (glucosyl)- glucoside	TLC- bioautograp hy	Sahu et al. 2012
Ficus coronata	Antibacterial activity against methicillin-resistant- <i>Staphylococcus aureus</i> and <i>Micrococcus luteus</i>	Skimmianine, 7- hydroxycoumarin, dihydrocoumarin, bergapten, chalepin, rutamarin, suberenol	HPLC-ESI- MSn and ESI-MSn	Smyth et al. 2012
Marine-derived <i>Streptomyces</i> sp. culture	Cytotoxicity	One unknown and one known compounds	microprobe NMR, ESI- MS	Mahyudin et al. 2012
Elaeocarpus chinensis	Cytotoxicity against a human colon cancer cell line	Epoxycucurbitacin derivatives and cucurbitacins	LC/MS	Pan et al. 2012
Seven most active among 289 fungal extracts	<i>In vitro</i> bioactivity towards leukemia cells	Ophiobolins	LC/DAD/ MS, explorative SPE	Bladt et al. 2012, Bladt et al. 2013

Tanacetum	In vitro membrane	Corvdaline.	NMR.	Konczol et
parthenium. Vinca	permeability assay for the	Vincamaiine, 11,13-	LC/MS	al. 2013
maior. Salvia	blood brain barrier	dihvdroparthenolide.		
officinalis, and		tetrahydropalmatine,		
Corvdalis cava		majdine, parthenolide,		
, ,		methyl carnosate.		
		epiisorosmanol		
Aglaia perviridis	Cytotoxicity against a human	Eight new compounds.	LC/MS	Pan et al.
0	colon cancer cell line	and 16 known		2013
		compounds		
Vulcanodinium	Cytotoxicity test against	Nakijiguinone A. N-	MS	Geiger et al.
rugosum	Neuro2A and KB cell lines	carboxy-methyl-		2013
(dinoflagellate)		smenospongine and		
		stachybotrin A		
<i>Carteriospongia</i> sp.	Antiproliferative activity	Homoscalarane-	UV, IR,	Harinantenai
and another similar	against three cancer cell lines	derived sesterterpenes	NMR	na et al. 2013
sponge		1		
Lipophilic extracts	<i>In vitro</i> anti-malaria and	Polyketides, lipids,	LC/MS,	Calcul et al.
from Chinese	cytotoxicity	diaporthochromones,	NMR	2013
mangrove endophytes		mycotoxins lipids		
Culture extract of	<i>In vitro</i> antimicrobial activity	Benzoic acid	TLC,	Rakshith et
<i>Xylaria</i> sp., an	against human and	derivative	LC/DAD/	al. 2013
endophytic fungus	phytopathogenic bacteria and		MS	
from Ficus pumila	fungi			
Premna odorata leaves	<i>In vitro</i> antimycobacterial test	1-heneicosyl formate,	GC/MS	Lirio et al.
		β-sitosterol,		2014
		stigmasterol and		
		diosmetin		
Three active plant	In vitro anti-fungal activity	Betulinic acid	MS, NMR	Bertrand et
extracts from French	against Candida albicans			al. 2014
Polynesia including				
Alphitonia zizyphoides				
Premna odorata	In vitro antimycobacterial	1-heneicosyl formate,	GC/MS,	Lirio et al.
Blanco leaves (from	activity	p-sitosterol,	NMR	2014
Philippines)		stigmasterol and		
		diosmetin		
Eriodictyon	In vitro ependymoma cell line	Flavonoids and	LC/MS/EL	Yang et al.
angustifolium and	bioassay	diterpenoids	SD/DAD	2014
Thuja occidentalis			database	
Piptocoma antillana	In vitro antiproliferative test	Two novel	MS, NMR	Liu et al.
leaves and twigs	against ovarian cancer cells	goyazensolide-type		2014
	and antiparasitic assay against	and two known		
	Plasmodium falciparum	sesquiterpene lactones		
Trigonella foenum-	In vitro SIRT6 (histone	Orientin and seventeen	LC/MS	Singh et al.
graecum seed extract	deacetylase involved in age-	other compounds		2014
	associated metabolic			
	disorders) binding test			

#### Dereplication procedures directly applied to crude extract collections (DEREP3)

The ability to directly identify molecular structures within crude extracts of natural products has become a critical step, either to cope with potent synergistic effects between constituents during biological evaluation, or even just to ensure that the material under examination presents an interest before going further with fractionation or purification steps. Here again, at the earliest stage of natural resource analysis, dereplication strategies have proved their value. It is obvious that the quick identification of known compounds directly in crude extracts is very ambitious because of the highly complex mixtures of non-fractionated samples. However, with the recent analytical advances, mainly in terms of resolution improvement and hyphenation capabilities, a range of efficient analytical systems are available today for the detailed analysis of complex mixtures (Funari et al. 2013, Michel et al. 2013, Carter 2014, Halabalaki et al. 2014). We will note as "DEREP3" all studies (n=50, 16 % of the total examined articles) found in the literature survey in which natural product dereplication consisted of the direct analysis of crude extract collections, either in combination with bioassays or not, but in all cases without fractionation or isolation of individual constituents. These studies are summarized in Table 4 and their workflow is illustrated in Fig. 4.



**Fig. 4.** DEREP3: a dereplication approach fully embedded in untargeted metabolomic analyses of natural extract collections

It emerged from the literature survey that different objectives could be reached by means of DEREP3-based workflows. Many of them aimed at the chemical profiling of crude extract collections by using pattern recognition tools or statistical treatments in an attempt to facilitate the visualization of either known or unusual spectroscopic features. In this way a substantial amount of time can be saved by focusing only on novel substances. It was reported that a hierarchical clustering analysis of GC/MS data obtained from 500 bacterial isolates allowed the selection of samples with a high probability of containing unknown natural products while avoiding the unnecessary analysis of samples of similar composition (Boroczky et al. 2006). In another interesting study, a collection of 30 New Zealand algae extracts were analyzed by a 2D NMR method based on the addition of HSQC spectra to construct a "stacked HSQC digital mask" displaying strong similarities between spectra as intense peaks (Popplewell and Northcote 2009). In order to enhance the unique signals present, implying potentially novel compounds, the mask was substracted from each individual HSQC spectrum. In this way a new bromophenol, colensolide A, was isolated from the red algae *Osmundaria colensoi* together with the known lanosol (2,3-dibromo-4,5-dihydroxybenzyl alcohol) and four of its derivatives.

A range of untargeted metabolomic studies fit perfectly with DEREP3 strategies and provide interesting perspectives especially in the chemical profiling of underexplored marine or microbial extracts. Investigation of marine species and microbes are indeed developing at an exponential rate, mainly based on phylogenetic analyses for drug lead discovery, but the vast array of available species and strains together with their unusual metabolite profiles as compared to plant species make it difficult to evaluate their metabolite composition (Rocha-Martin et al. 2014). It was reported in 2012 that the investigation of filamentous fungi for anticancer drug leads resulted in the isolation and characterization of only 140 compounds over the four previous years, and that only 30% of these represented new chemical entities (El-Elimat et al. 2012). Dereplication of microbial culture collections using DEREP3-based metabolomic procedures can therefore be helpful to classify strain collections *via* a chemotyping approach. Moreover, DEREP3-based strategies should accelerate the evaluation of metabolite diversity, mainly by avoiding redundancy in the identification of metabolites produced by the species under examination and by maximizing microbial natural product libraries that can be generated from collections of microorganisms (Berdy 2005, Larsen et al. 2005, Silver 2006, Liu et al. 2012, Tawfike 2013, Ito and Masubuchi 2014). The direct analysis of fungi directly in fermentation extracts using LC/MS, pattern recognition tools and databases was reported in several papers as an efficient alternative to discover novel molecules with unknown activities (Martin et al. 2012, El-Elimat et al. 2013). In another example, a metabolomic approach combining HR-FTMS and NMR spectral data to PCA, HCA and OPLS-DA statistical treatments was used for the dereplication of metabolites in antitrypanosomally active spongeassociated bacterium Actinokineospora sp. obtained from four different fermentation conditions, the main objective being to identify the best culture one-strain-many-compounds condition for the isolation of novel bioactive metabolites (Abdelmohsen 2014).

Other interesting studies performing "co-cultures" of microorganisms to enhance the diversity of metabolite production *via* the activation of silent genes have followed a DEREP3 workflow to rapidly determine the presence of novel compounds while leaving aside those already known (Butler et al. 2012, Bertrand et al. 2013). For example, ten known compounds were identified by dereplication in a crude extract of the sponge-derived actinomycetes *Actinokineospora* sp. while three unexpected natural products not detected in the single culture were identified when *Actinokineospora* sp. was grown in co-culture with *Nocardiopsis* sp. (Dashti, Grkvic et al. 2014).

Another interesting aspect of DEREP3 strategies arises from the possibility to directly correlate biological properties to chromatographic and spectroscopic features of the crude analyzed samples. In this case, the purpose is to link the chemical characteristics of natural compounds to a specific biological activity. With the help of the ChemGPS-natural product database, it was shown that the inhibition of cyclooxygenase enzymes involved in inflammatory processes is frequently correlated with the presence of at least one ring in the metabolite structure. Fragments exhibit structural rigidity, and compounds have a relatively large molecular volume (Larsson et al. 2005). In another example, a screening technology combining on-line biological analysis with the resolution of LC/MS for structure elucidation was developed to directly analyze estrogen-receptor binding compounds in complex mixtures of a large plant extract library (Schobel et al. 2001). LC/MS signals of compounds from 22 phenolic extracts of extra virgin olive oil showing an activity against a breast cancer cell line were also identified by applying a correlation analysis between particular spectral features and the observed activity. From this method a model was estimated to predict the activity of new samples (Roldan et al. 2013). Optimization of such strategies in future research efforts could significantly accelerate the discovery of novel active substances and help our understanding of natural products with respect to structure-activity relationships.

**Table 4.** Dereplication strategies published from 1990 to 2014 using a DEREP3 workflow(from oldest to most recent)

Screening of a collection of 116 marine sponges, ascidians, and cnidarians for their antifungal activity against Candida albicansExtract prioritization(Antonio and Molinski 1993)Screening of a collection of aqueous extracts of terrestrial plants, cyanobacteria, and marine invertebrates for their AIDS-antiviral activityExtract prioritization(Cardellina et al. 1993)Screening of 38,000 extracts from plants and microorganisms for antibacterial activity. Four promising extracts were retained after chemical dereplicationEarly identification and elimination of known compounds(Ramakrishn a et al. 1999) compoundsLC/MS analysis of a plant extract library combined on-line to an estrogen receptor binding assayAcceleration of bioactive compound characterization(Schobel et al. 2001)Curie-point pyrolysis MS and numerical taxonomic analysis of 44 deep-sea Rhodococci, clustering according to PyMS fingerprints and comparison with conventional microbialTaxonomic classification(Colquhoun et al. 2000)
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Single quadrupole LC/MS discrimination of novel from known Early identification and (Gilbert et al.
compounds in crude biologically active extracts elimination of known 2003)
Compounds
Screening of lungal metabolites and mycoloxins in cultured Early identification and (Nielsen and
extracts using a LC/U V/MS micro-scale method and a MS- librory
Indrary     Compounds     2003)       Samaring of 16 Durivillium analise groups in different matrings     Englustion of strain shamical     (Nisland strain shamical)
Screening of 16 <i>Peniculuum</i> species grown in different matrixes, Evaluation of strain chemical (Neisen et al. 2004)
and evaluation of produced metabolites <i>via</i> a chemo-diversity diversity al. 2004)
Here of the ChamCDS database for a set of natural products Investigation of correlations(Loreson et
out the Chemory database for a set of natural products investigation of correlations (Larsson et avkibiting avalages a and/or 2 (COX 1/2) inhibition
exhibiting cyclooxygenase-1 and/of -2 (COX-1/2) minorition between structural features al. 2003)
Screening of 456 bacterial isolates from marine sponges by Papid selection of stroins (Diackmann)
using Integet Coll MALDI TOF MS and protoometric clustering representing representi
of the strains into 11 groups corresponding to particular species subsequent chemical
of the strains into 11 groups corresponding to particular species subsequent chemical
Analysis of four different collections of the sponge genus Zyzzy/ Search for anticancer (Dijoux et al.
Analysis of four different conections of the sponge genus $Zy_{z,yu}$ Search for anticancer (Dijoux et al.
Cancer Institute's 60 human tumor cell lines
Analysis of 500 bacterial isolates by GC/MS combined to UCA Early identification of known (Percentry et
Analysis of 500 bacterial isolates by GC/1415 combined to TICA Early identification of Known (B010c2ky et compounds al 2006)

Development of a simple method for high-throughput	Removal of promiscuous	(Appleton et
prefractionation of crude extracts	inhibitors and interference	al. 2007)
	compounds during	
	enzyme/protein-based assays	
MALDI-TOF-imaging of intact marine cyanobacteria (Lyngbya	Characterization of the	(Esquenazi et
majuscula 3L and JHB, Oscillatoria nigro-viridis, Lyngbya	spatial distribution of natural	al. 2008)
bouillonii, and Phormidium species) and sponges	products	
Analysis of myxobacterial metabolites in nine Myxococcus	Investigation of potentially	(Krug et al.
species using LC/ESI-TOF combined to PCA	novel natural products,	2008)
	prioritization of candidates	
Chemical profiling of fungal or bacterial extracts using an HPLC	Demonstration of the	(Lang et al.
bioactivity profiling/microtiter plate technique in conjunction	discriminating power of H-1	2008)
with capillary probe NMR instrumentation and databases	NMR as a dereplication tool	
Rapid determination of log P using UHPLC profiling gradients	Implementation of	(Eugster et
on a representative library of natural products and investigation	dereplication database for	al. 2009)
of the relations log P-log k at different pH and UHPLC	crude plant extract profiling	
conditions	studies	
Monitoring of different substrate utilization, growth, secondary	Microbial drug discovery,	(Singh 2009)
metabolite and antimicrobial profiles of some common	dereplication of fungi and	
filamentous fungal cultures using the Biolog FF MicroPlate,	differentiation of closely	
LC/MS and antimicrobial assays	related variants within one	
	species	
Screening of aqueous and ethanol extracts of 5 South African	Removal of non-specific	(Klos et al.
medicinal plants for activity against HIV	undesired tannins and	2009)
	polysaccharides	
NMR analysis of 30 New Zealand marine algal extracts, addition	Extract prioritization	(Popplewell
of all HSQC spectra to construct a "stacked HSQC digital mask"		and
and substraction of this mask from each individual HSQC		Northcote
spectrum to enhance the unique signals		2009)
Metabolomic analysis of crude extracts obtained from	Identification of novel	(Crevelin et
actinomycetes in differents cultivation medium by direct	microbial metabolites	al. 2010)
infusion MS and LC/MS		un <u>2010</u> )
Monitoring of aerial spore mass, pigment colours formed on	Bioprospection and	(Antony-
oatmeal agar and capacity to produce melanin pigments of a	ecological study	Babu et al.
range of alkaliphilic <i>Streptomycetes</i> isolated from a beach and	······	2010)
dune sand system. Computer-assisted method for colour-		/
grouping and comparison to PCR-data		
LC/DAD/MS and NMR analysis of active crude extracts	Understanding molecular	(Castro-
from Abarema lusoria and Calea pinnatifida and comparison	relationships on dynamic	Gamboa et
with <i>in silico</i> databases	natural matrixes	al. 2010)
Explorative solid-phase extraction and ion-exchange	Extract prioritization.	(Mansson et
chromatography for the chemical profiling of microbial extracts	mapping of biological	al. 2010)
	activities	
Chemical screening of thirteen antibiotics and four cultivation	Identification of novel potent	(Kamenik et
broths by LC/DAD and fingerprint analysis based on polarity,	antibiotics among microbial	al. 2010)
UV spectra and acid-base properties	metabolites	

High-throughput LC/MS analysis of 16 025 microbial extracts,	Investigation of extract	(Ito et al.
binning by nominal mass and retention time, data visualization	chemical diversity	2011)
by scattered plots and elimination of the ubiquitous peaks to		
focus on unique compounds		
Analysis of 249 unidentified bacterial isolates retrieved from the	Comparison of the	(Ghyselinck
rhizosphere of potato plants using MALDI-TOF MS, repetitive	taxonomic resolution	et al. 2011)
PCR or 16S rRNA gene sequence analysis and cluster analysis	obtained by different	
of the profiles obtained from the different techniques	techniques for a broad	
	diversity of bacteria	
Analysis of fungal secondary metabolites in culture extracts	Search for novel potent	(El-Elimat et
using LC/DAD/MS with MS/MS spectral database	anticancer compounds before	al. 2012)
	engaging in isolation process	
Metabolomic analysis of marine-derived bacterial natural	Strain prioritization in a drug	(Hou et al.
products using LC/MS with PCA	discovery program	2012)
Molecular virtual design using NMR analysis of highly active	Exploring Brazilian	(Castro-
crude extracts from Cerrado and Atlantic Rainforest specimens	biodiversity	Gamboa et
and endophytic fungi and microorganisms derived from specific		al. 2012)
rizhosphere habitats		
Chemical profiling of <i>Thymus vulgaris</i> extracts using	Investigation of extract	(Pieri et al.
LC/SPE/NMR and PCA to determine the discriminating	chemical diversity	2009, Pieri et
constituents		al. 2012)
Chemical profiling of 15 extracts of various organs of six <i>Lippia</i>	Early identification of known	(Fugster et
species using LC/DAD/TOE-MS_MS spectral databases and	compounds	(Lugster et)
HCA to highlight discriminating metabolites among species	compounds	Funari et al
The A to might discriminating inclabolities alloing species		2012
		2012)
Chemical profiling of 15 extracts of six Lippia species by	Early identification of known	(Funari et al.
LC/DAD, comparison with standard compounds and PCA	compounds	2012)
Investigation of eight marine-derived <i>Furcatum</i> and <i>Penicillium</i>	Identification of novel potent	(Vansteeland
strains grown on six different culture media using LC/DAD/MS <sup>n</sup>	bioactive compounds	t et al. 2012)
Screening of non bioactive fungal extracts fermented in 8-	Discovery of novel	(Martin et al.
medium nutritional arrays using LC/MS combined to an in-	molecules with unknown	2012)
house database	activities	
Chemical profiling of 22 phenolic extracts of extra virgin olive	Rapid identification of	(Roldan et al.
oil using LC/MS and <i>in vitro</i> cytotoxicity bioassay. Correlation	bioactive compounds directly	2013)
analysis between the activity and LC/MS data and development	within crude extracts	
of a model to predict the activity with further new samples		
Metabolomic analysis of Streptomyces isolated from	Discrimination of microbial	(Forner et al.
geographically varied environments using LC/MS combined to	strains and identification of	2013)
bucketing and presence-absence standardization strategy, PCA	novel compounds	
and HCA		
Dereplication of <i>Penicillium</i> strain extracts grown on various	Assessment of toxigenic	(Geiger et al.
media using HPLC-UV/DAD-MS/MS	risks associated to fungal	2013)
	strains	
LC-MS-based metabolomic analysis of pure strain cultures and	Identification of novel	(Bertrand et
--	-------------------------------	---------------
co-cultures of Trichophyton rubrum and Bionectria ochroleuca	metabolites induced by	al. 2013)
	fungal interaction	
Metabolomic analysis of 600 fungal co-cultures in solid media	Identification of novel	(Bertrand et
using UHPLC-TOF-MS, statistical comparison with the	metabolites induced by	al. 2013)
metabolite profiles obtained from mono-culture	fungal interaction	
Screening of filamentous fungi cultured extracts for the	Early elimination of extracts	(El-Elimat et
discovery of anticancer drug using LC/DAD/MS and MS/MS	containing known	al. 2013)
	compounds	
Chemical profiling of 22 African propolis samples using	Rapid identification of	(Zhang et al.
LC/UV/ELSD, LC/MS, GC/MS, LC/DAD/MS <sup>2</sup> , heat mapping	discriminating metabolites	2014)
LC-UV and ELSD data, PCA and exact mass search in the DNP		
Morphological and chemical characterization of Fusarium	Chemical investigation of	(Short et al.
keratoplasticum sp. and Fusarium petroliphilum	Fusarium species	2013)
Characterization of the pigment profiles of 400 bacterial isolates	Taxa discrimination and	(Stafsnes et
using MALDI-TOF-MS dendrograms	identification of novel	al. 2013)
	carotenoids with UVA-Blue	
	light absorbing properties	
Screening of a library of 3120 natural extracts with potent	Biological screening of novel	(Wong et al.
antibiotic properties, clustering of bioactivity profiles	potent antibiotics	2013)
Metabolomic analysis of 278 extracts from Malaysian	Prioritization of extracts	(Samat et al.
biodiversity using LC/MS combined to PCA.	potentially containing novel	2014)
	photosensitizers	
Metabolomic analysis of antitrypanosomally active sponge-	Identification of the best	(Abdelmohse
associated Actinokineospora sp. obtained in four fermentation	culture one-strain-many-	n 2014)
conditions using HR-FT-MS and NMR combined to PCA, HCA	compounds conditions	
and OPLS-DA		
Chemical profiling of 77 bacterial extracts isolated from cold	Acceleration of strain	(Macintyre et
water marine invertebrates from Scotland using LC/MS	prioritization	al. 2014)
combined to PCA and NMR ( <sup>1</sup> H and COSY)		
Dereplication of Aspergillus carbonarius and Penicillium	Screening of novel bioactive	(Klitgaard et
melanoconidium extracts using LC/DAD/ESI-MS, overlay of	metabolites	al. 2014)
automatically generated extracted-ion chromatograms to		
visualize novel peaks		

Dereplication and targeted chemical profiling: two terms coming down to the same

approach (DEREP4)

As mentioned above, in a targeted metabolite profiling approach a predefined number of compounds or a particular chemical class of compounds is investigated and the molecular structures are identified. Remembering that the literal definition of dereplication is "*the rapid* 

*identification of known compounds*", and assuming that the range of compounds investigated in targeted chemical profiling studies can also be known, it can be considered that both concepts are quite similar in many cases and that a targeted chemical profiling is necessarily a kind of dereplication, even if in contrast dereplication does not always correspond to a targeted chemical profiling approach.

In the literature survey, 57 studies (18 % of the total examined articles) involving dereplication were used a targeted chemical profiling approach. These studies are summarized in Table 5 and are noted as DEREP4 strategies. Their global workflow is given in Fig. 5.



**Fig. 5.** DEREP4: a targeted dereplication workflow focused on the identification of a predefined group of known molecules

The main objective of DEREP4-based studies is either to analyze a particular group of molecules presenting a biological interest or on the contrary to identify potentially toxic or undesired constituents of crude natural extracts. For example, the dereplication of free sugars and polyols was performed by GC/MS in a study investigating novel natural sweeteners. Among the sweet-tasting species analysed, the extracts containing high levels of these

compounds were regarded as lower priority leads and thus directly removed from further consideration, enabling a significant time gain (Chung et al. 1997). DEREP4 approaches are also commonly used to assess the quality of botanical drugs that must be checked for authentication, contamination, and identification of active substances or safety validation. Such procedures are based on the fact that all natural extracts, although highly complex, contain secondary metabolites which are specific to the species under study. During analytical data capture, the set of signals corresponding to these specific metabolites becomes a chemical fingerprint and the most characteristic signals are called "chemical markers". For instance three major concentrated stilbenoids quickly identified by a HPTLC/MS-based dereplication approach and exhibiting skin-anti-ageing properties were reported as relevant markers of a range of orchid species (Williams et al. 2012). Of course, concentration variability of these chemical markers in natural raw materials is inevitable due to inherent factors such as genotype variability, growing conditions, and climate or soil type. By tracking chemical markers over different steps of the manufacturing processes, the DEREP4 approach can assess this variability and minimize the potential composition heterogeneity of the final ingredient from one batch to another. This procedure forms an integral part of the standardization of botanical drugs, a major task which is necessary to adjust the concentration of the active substances and ensure a reproducible quality of the final products.

In the vast majority of targeted chemical profiling studies, natural metabolites were analyzed by mass spectrometry. This can be explained by the fact that a group of targeted compounds generally comprises structurally related molecules, consequently exhibiting very similar fragmentation patterns. The use of MS-based libraries containing fragmentation spectra and substructure data of the targeted compounds is thus very useful in the case of such dereplication approaches. For instance, a range of taxoids were successfully identified in a crude extract of *Taxus wallichiana* with the help of a MS library containing fragmentation spectra of 3 taxoid standards and substructure spectra of 139 previously reported toxoids (Stefanowicz et al. 2001). Similarly, it was found that retro-Diels-Alder rearrangement occurring during the MS fragmentation of a xanthone skeleton produce characteristic fragment ions which could be used to target prenylated xanthones in plant extracts such as *Garcinia* plants (Zhou et al. 2008). The fragmentation pattern of flavonoids and their glycoside derivatives is also commonly used to dereplicate this class of compounds in crude natural extracts through targeted chemical profiling approaches (Constant et al. 1997, Waridel et al. 2001, Tchoumtchoua et al. 2013). Mass accuracy and isotopic patterns are other data recovered from MS and MS<sup>n</sup> acquisition that can be used to build up libraries of natural metabolites for dereplication purposes (Gomez-Romero et al. 2011).

It can be supposed that NMR-based metabolite profiling strategies are less efficient to perform the targeted chemical profiling of a predefined set of compounds, mainly because of the strong NMR signal overlaps observed when analysing a group of structurally related molecules. However, several studies have demonstrated that this issue can be overcome by focusing on specific NMR signals of the targeted molecules. The screening of several brominated tyrosine derivatives named bastadins in a crude extract of the marine sponge *Lanthella basta* was performed for instance by <sup>1</sup>H NMR detection of their characteristic methoxy signals (Franklin et al. 1996). In the same way, diagnostic <sup>1</sup>H NMR signals of epoxide and conjugated diene moieties have been used to dereplicate macrocyclic trichothecenes in filamentous fungal species in the context of new anticancer lead discovery (Sy-Cordero et al. 2010).

Finally, it should be mentioned that targeted chemical profiling approaches not only enable the identification of previously known compounds, but also promote the discovery of additional novel metabolites, often corresponding to structural analogs of the predefined class of targeted compounds.

**Table 5.** Dereplication strategies published from 1990 to 2014 using a DEREP4 workflow(from oldest to most recent)

Targeted compounds and natural sources	Analytical technique(s)	Ref
Cucurbitacins in Iberis amara, Begonia plebeja and	Computer-Assisted	(Fuller et al. 1994)
Gonystylus keithii	recognition of cytotoxicity	
	profiles	
Elaiophylin and geldanamycin in a range of microbial	CPC/DAD, LC-MS	(Alvi et al. 1995)
broths		
Two known antibiotics teicoplanin and phenelfamycin	LC/UV/ESI-MS/MALDI-	(Ackermann et al.
in several microbial crude extracts	MS	1996)
Bastadins from Lanthella basta	<sup>1</sup> H NMR	(Franklin et al.
		1996)
Free sugars and polyols in six sweet-tasting plant	GC/MS	(Chung et al.
extracts		1997)
Flavonoids and flavonoid glycosides using an aqueous	LC/MS and CID for	(Constant and
extract of Eugenia jambos as a model	fragmentation patterns	Beecher 1995,
		Constant et al.
		1997)
Potentially interfering polyphenols during the biological	LC/UV/MS	(Kinghorn et al.
screening on in vitro cytotoxicity bioassays of 3000		1995, Cordell and
plant extracts from tropical rainforests against a panel		Shin 1999,
of human cancer cell lines.		Kinghorn et al.
		1999)
Seven destruxing in the fungus <i>Metarrhizium anisopliae</i>	HPLC/ESI-TOF-MS	(Potterat et al.
		2000)
Taxoids from Taxus wallichiana	MS <sup>n</sup>	(Stefanowicz et al.
		2001)
C-glycosidic flavonoids	LC/MS/MS using Q-TOF	(Waridel et al.
	and IT under various CID	2001)
	energy conditions	
Sesquiterpenes from two Indo Pacific sponges	NMR	(Stessman et al.
Lendenfeldia frondosa and Hyrtios sp.		2002)
Trichothecenes from seven microbial Trichoderma	LC/UV/vis and ESI-MS	(Nielsen et al.
species grown on several solid and liquid media		2005)
Betaxanthins from yellow-orange cactus pear fruits and	<sup>13</sup> C NMR	(Stintzing et al.
yellow Swiss chard petioles		2006)
High molecular weight polysaccharides from several	DOSY-based NMR	(Politi et al. 2006)
mushroom extracts		
Pentacyclic triterpenoids as their methyl esters	GC/ESI-MS	(Gu et al. 2006)
(oleanolic acid, betulinic acid and ursolic acid) with		
potent anti-tubercular activity		
Saponins and acyclic sesquiterpene oligoglycosides in	LC/UV/ESI-MS, MS/MS	(Murgu and
the fruits of Sapindus saponaria		Rodrigues 2006)
Anti-inflammatory compound vicenin-2 in Leucopogon	LC/ESI-MS <sup>n</sup>	(de Moraes et al.
ericoides		2007)

(+)-psymberin and a new brominated cyclic peptide (-)-	ESI-MS, NMR	(Robinson et al.
psymbamide A from the sponge Psammocinia aff.		2007)
bulbosa		
Novel phenolic compactin analogue from Penicillium	UV-guided strategy, NMR,	(Larsen et al.
solitum	MS	2007)
Valerenic acid in a valerian extract and other GABA(A)	HPLC	(Kim et al. 2008)
receptor ligands in plant and fungal extracts		
Streptothricin-like compounds in the fermentation broth	Ion-pair LC/ESI-MS	(Ji et al. 2008)
of Streptomyces qinlingensis		
Polyprenylated xanthones in Garcinia	On-line LC/ESI-QTOF-	(Zhou et al. 2008)
xipshuanbannaensis	$MS^3$	
Alkaloids in several Colchicum species	LC/MS, LC/UV/DAD	(Alali et al. 2008, Gharaibeh et al.
Secoiridoid glycosides in Sarracenia alata	capillary-scale NMR, ESI- MS	(Hu et al. 2009)
Plumeran indole alkaloids in a stem bark extract of	ESI-MS/MS	(Aguiar et al.
Aspidosperma spniceanum		2010)
Bufadienolides in the TCM toad skin	UHPLC/ESI-Q-TOFMS	(Liu et al. 2010)
Bromotyrosine-derived metabolites in 14 specimens of	UV, LC/DAD/MS	(Silva et al. 2010)
Aplysina spp. marine sponges		
Heterocyclic nerolidol derivatives from the anti-	<sup>1</sup> H NMR with focus on	(Inui et al. 2010)
tuberculosis active fractions of the inner stem bark of	spin-spin coupling patterns	
Oplopanax horridus		
Macrocyclic trichothecenes from different fungal	UV spectra, <sup>1</sup> H NMR	(Sy-Cordero et al.
species		2010)
Bastadins from the marine sponge <i>Ianthella</i> reticulate	NMR	(Calcul et al. 2010)
Phenolics in propolis and lyophilisate of some	LC/ESI-QTOF	(Gomez-Romero
vegetables selected for their antioxidant properties		et al. 2011)
Stilbenoids in several species of the Orchidaceae family	MicroCryoProbe NMR	(Williams et al. 2012)
Pregnane glycosides and genins from the TCM	HPLC/ESI-MS <sup>n</sup>	(McGarvey et al.
Marsdenia tenacissima		2012)
Aporphine and oxoaporphine alkaloids from Unonopsis	direct infusion in ESI-IT-	(da Silva et al.
guatterioides	MS	2012)
Active photosensitisers possessing a cyclic tetrapyrrole	LC-DAD-MS	(Tan et al. 2012)
in 15 plant extracts		
Nonribosomal peptides	MS/MS, chemoinformatic library	(Ibrahim et al. 2012)
Bisbenzylisoquinoline alkaloids from the fruit of	UV, MS	(Klausmeyer et al.
Gyrocarpus jacquinii		2012)
Annonaceous acetogenins from Annona muricata	LC/LTQ-Orbitrap	(Le Ven et al. 2012)
170 known triterpenes from Actaea species	<sup>1</sup> H NMR and predictive model using classification binary trees	(Qiu et al. 2012)

Aliphatic alkaloids from Huperzia selago and	LC/SPE/NMR	(Johansen et al.
bisbenzylisoquinoline alkaloids from Triclisa patens		2012).
Minor cytotoxic chlorinated valepotriates from whole	HPLC-PDA-MS, TLC,	(Lin et al. 2013)
plants of Valeriana jatamansi	NMR	
Steroidal alkaloids in complex plant extracts of <i>Buxus</i>	ESI-TOF-MS/MS	(Musharraf et al.
species		2013)
Thiazolyl peptides from	Miniaturized 96-well SPE,	(Singh et al. 2013)
	HR-FT LC/MS and library	
Lasiodiplodins from a cytotoxic extract obtained from	ESI-MS, NMR	(Sultan et al. 2014)
<i>Lasiodiplodia theobromae</i> , an endophyte from the root		
tissues of Mapania kurzii		
Ubiquitous flavonoids and related plant constituents	Q-TOF, IT and MS <sup>2</sup>	(Wolfender et al.
	library	2000)
Rhizoxins and rhizonins in different Zygomycetes	LC/DAD/HR-MS	(Jennessen et al.
grown on a range of semisynthetic and natural		2005)
substrates		
Stilbenoids in wine fractions	LC/DAD/MS, LC/NMR	(Pawlus et al.
		2013)
Flavonoids and benzophenone derivatives from Qualea	HPLC/ESI-QTOF-MS/MS	(Neto et al. 2013)
grandiflora and Qualea cordata extracts		
Isoflavonoids from a stem bark of Amphimas	UHPLC/MS, MS/MS	(Tchoumtchoua et
pterocarpoides		al. 2013)
Pyrrolamide compounds including congocidine and	genome scanning and	(Hao et al. 2014)
distamycin in fermentation culture of <i>Streptomyces</i>	precursor ion scan MS	
netropsis		
Triterpenes from various Actaea species	2D-NMR barcoding	(Qiu et al. 2014)
Trace-levels of acetogenins in an Annona cherimolia	LC/HRMS with	(Le Ven et al.
fruit-based alcoholic beverage	postcolumn infusion of	2014)
	lithium iodide	
Surfactin-type lipopeptides in <i>Bacillus</i> cultures from	LC/HRMS	(Juola et al. 2014)
commercial Japanese foods		

## Molecular genetics and taxonomic dereplication of microbial strains (DEREP5)

Microbial natural products are a major resource for the development of novel drugs in pharmaceutical industry. However, the chemical characterization of microbial secondary metabolites is currently hampered by the laborious isolation processes when working with culture broths, and by the limited availability of structural and spectral libraries for microbial metabolites. In addition, microbial culture collections are extremely diverse, and thus a major challenge in their rational exploitation is the reduction of unnecessary or redundant testing of strains. In this sense, a range of dereplication tools mainly based on phenotypic characteristics or genetic differences have been developed to select microbial isolates at the species level. Genetics and metabolic engineering are currently an integral part of these dereplication strategies. In recent years, the exploration of new taxa or ecological niches through metagenomic approaches have provided access to a large number of gene sequences that can now be investigated not only in well-studied bacteria but also in microorganisms that were previously neglected as natural ressources. The metagenome can be defined as a "collective genome" including isolated microbial DNA of all microorganisms present in a particular habitat (Handelsman et al. 1998). "DEREP5" will thus denote all studies (n=22, 7 % of the total examined articles) found in the literature survey that involved a genetic approach for the dereplication of microbial strains. These studies are all included in Table 6 and their workflow is illustrated in Fig. 6.



Fig. 6. DEREP5: A taxonomic dereplication workflow based on gene-sequence analyses of collection of microbial strains

It must be specified that many DEREP5 studies were performed on microbial strains or isolates "as a whole" and did not systematically include a detailed analysis of the metabolites produced by the selected microorganisms. Nevertheless, such approaches have already led to the construction of extensive taxonomy-guided microbial natural product libraries, and thus enables for instance biodiversity surveys, strain prioritization, spatial distribution characterization, biosynthetic pathway elucidation, discovery of untapped sources of natural products, and detection of uncommon genera or species among microbes. Most of the current protocols for the taxonomic dereplication of microbial strains involve 16S rDNA gene sequencing or PCRfingerprinting of large collections of isolates followed by clustering analysis in order to correlate phylogenic similarities or to highlight discriminating strains. Once microorganisms have been taxonomically identified and revealed as promising leads, more sensitive analytical tools can be implemented to investigate metabolite profiles and potentially identify novel bioactive compounds. Strategies for in silico-guided identification of secondary metabolites by genome mining have been highlighted in several reviews (Zerikly 2009, Helfrich et al. 2014, Helfrich 2014). Such strategies seem promising, but although novel compounds have been revealed in activity-driven screening of gene libraries, the frequency of the identified compounds is relatively low (Novakova and Farkasovsky 2013). Research efforts are currently being reinforced to better understand microbial and chemical ecology, improve microbial culture conditions, high-throughput cultivation and performance of large-scale biological screening assays (Ashforth et al. 2010, Baltz 2010, Penesyan et al. 2010). Improvements in sequencing capabilities would also contribute to this area, particularly regarding primer design, increase of insert sizes and diversification of library hosts (alternative to Escherichia coli-based libraries).

It must finally be mentioned that the large majority of DEREP5 studies are aimed at screening novel antibacterial metabolites. Pathogen resistance to currently available antibiotics has indeed

reached a very critical level. Over the last twenty years, most pharmaceutical companies have abandoned natural product chemical profiling investments in favor of large and cost-effective libraries of synthetic molecules. However, it is obvious that further improvements in genome sequencing technologies, together with the use of modern analytical systems, computer tools and application of reliable and pertinent biological assays will progressively solve the speedlimiting dereplication obstacles and lead these companies to refocus on natural product candidates (Kirst 2013). Several reviews addressing the value and uses of molecular geneticsbased dereplication procedures applied to natural microbial metabolites are available in the literature (Strobel 2002, Knight et al. 2003, Leeds et al. 2006, Liu et al. 2010, Genilloud et al. 2011, Roemer 2011, Genilloud 2014, Sandiford 2014).

**Table 6.** Molecular genetic-based strategies developed for the taxonomic dereplication of microbial natural products (DEREP5)

Strategy	General goal	Ref
Investigation of unusual strains of actinomycetes and	Detection of uncommon	(Donadio et al.
filamentous fungi by reconstruction of gene clusters from	genera of soil microbes	2002)
small segment of cloned DNA and preparation of large-		
insert libraries		
Identification of 26 closely related Streptomyces soil	Discrimination between	(Ritacco et al.
isolates using rDNA sequencing, MIDI fatty acid analysis,	morphologically similar	2003)
and LC-MS profiling of fermentation extract	strains	
Screening of 20 Micromonospora isolates from UK coastal	Investigation of strain	(Zhao et al.
sediments, taxonomic classification by rDNA sequencing,	diversity	2004)
fluorescent amplified fragment length polymorphism		
(AFLP), and Fourier transform infrared spectroscopy (FT-		
IR)		
Targeted analysis of polyketide synthases and	Characterization approach	(Ayuso et al.
nonribosomal peptide synthetases amplified sequences in a	for Actinomycetes	2005)
collection of wild-type Actinomycetes isolated from		
tropical soil samples evaluated for the production of		
antimicrobial activities		
Dereplication of microorganisms collected in 12	Investigation of biodiversity	(Huys et al.
aquaculture sites in Southeast Asian and using PCR	and environmental	2007)
fingerprinting	distribution of	
	chloramphenicol-resistant	
	(CmR) mesophilic	
	heterotrophs	

Evaluation of the antimicrobial activity and sub-grouping	Investigation of	(Hakvag et al.
of 217 streptomycetes isolates from the water surface	Actinomycetes from the	2008)
microlayer in Norway by phylogenetic analysis and 16S	water surface microlayer as	
rDNAs sequencing. 7 isolates with identical 16S rDNA	a source of new	
sequences were further studied for the presence of PKS	antimicrobial agents	
type I genes		
RNase P RNA gene (rnpB) sequencing of 50 myxobacteria	Development of RNase P	(Monciardini et
strains for the faster and cheaper discrimination of similar	RNA gene (rnpB)	al. 2008)
strains as compared to 16S rDNA sequencing	sequencing as a tool for	,
	molecular dereplication	
Isolation of <i>Micromonospora</i> strains on the basis of typical	Taxonomical classification	(Maldonado et
colonial and pigmentation features and 16S rRNA gene		al. 2008)
sequencing analyses and correlation with molecular		,
fingerprinting		
Dereplication of 127 <i>lactobacilli</i> isolates by (GTG)(5)-	Investigation of lactic acid	(Solieri et al.
PCR fingerprinting, phylogenetic analysis of the 51	bacteria strain diversity in	2012)
resulting genotypes using a combined amplified 16S rDNA	31 ripened Parmigiano	,
restriction analysis (16S-ARDRA), species-specific PCR	Reggiano cheeses	
assays and 16S rRNA gene sequencing		
ITS1 and ITS4-rDNA sequencing of four endophytic fungi	Investigation of new potent	(Hazalin et al.
extracts evaluated in parallel for their cytotoxic activity.	cvtotoxic fungal extracts	2012)
identification of known compounds using an UV library		
Multigene analysis, combined with phenotypic characters	Taxonomic classification of	(Houbraken et
and extrolite profile of <i>Penicillium</i> sect. <i>Chrysogena</i>	penicillin-producing strains	al. 2012)
species		,
Development of a web-based bioinformatics platform	Development of	(Yu et al. 2006)
(FastGroupII) to dereplicate large 16S rDNA libraries and	bioinformatic tools for the	
application on a set of 16S rDNA sequences from coral-	analysis of high-throughput	
associated bacteria	16S rDNA sequencing	
	databases	
Automated ribotyping of 157 isolates following restriction	Dereplication of a large	(Sheffield et al.
enzyme digestion, classification into 23 ribogroups	collection of phenotypically	2006)
representing a 85% reduction of the number of isolates in	ambiguous bacterial isolates	,
the library		
16S rRNA, gyrB and recA gene sequencing of thirteen <i>P</i> .	Selection of strains for	(Vynne et al.
<i>luteoviolacea</i> strains to determine whether chemotype and	antibiotic discovery	2012)
activity profile can be reflected by phylogenetic clustering	5	,
Dereplication of 433 Danish surface-ripened cheeses	Phylogenetic analysis of	(Gori et al.
bacterial isolates using (GTG)5-PCR fingerprinting, and	bacterial diversity in Danish	2013)
identification of 217 bacterial and 25 yeast isolates by 16S	cheeses	/
rRNA gene sequencing		
Characterization of microbial eukaryotic populations	Phylogenetic analysis of	(Garcia-
associated with Pyrenean glaciers using molecular 18S	microbial eukarvote	Descalzo et al.
rRNA-based approaches, amplifying community DNA and	diversity in icv regions	2013)
constructing clone libraries with 18S rRNA primers	, , , , , , , , , , , , , , , , , , ,	/
Taxonomic identification and phylogenetic reconstruction	Assessment of bacterial	(Stets et al.
of 129 collular extracts of unless accorded hostorial	isolate diversity	2013)

isolates using 16S rPNA game sequencing and whole call		
Isolates using 105 IKIVA gene sequencing and whole cen		
MALDI-TOF-MS analysis		
DNA expression array analysis of Aspergillus nidulans in	Elucidation of biosynthetic	(Andersen et al.
combination with legacy data to form a comprehensive	pathways of fungal	2013)
gene expression compendium and data clustering to	secondary metabolites	
identify cross-chemistry between physically separate gene		
clusters		
Development of a two-pronged approach for the	Rapid identification of the	(Orelle et al.
characterization of inhibitors of protein synthesis (ChIPS)	site and mode of action on	2013)
by engineering antibiotic-hypersensitive Escherichia coli	the bacterial ribosome	
strains that contain only one rRNA operon and elucidation		
of the mode of antibiotic action by monitoring drug-		
induced ribosome stalling on mRNA		
PCR analysis of 100 strains randomly selected from an	Extract prioritization by	(Xie et al.
actinomycete collection to evaluate their ability to	biosynthetic potential survey	2014)
biosynthesize aromatic polyketides, reduced polyketides,		
nonribosomal peptides, and diterpenoids		
Investigation of a collection of Aspergillus nomius strains	Characterization of	(Massi et al.
from Brazil nuts including morphological characters,	Aspergillus species in Brazil	2014)
RAPD and AFLP profiles, partial $\beta$ -tubulin and calmodulin	nuts	
nucleotide sequences, aflatoxin patterns, as well as		
tolerance to low water activity in cultured media		
Investigation of Streptomyces lavendulae phenotypic	Identification of potentially	(Sladic et al.
cluster for production of lipstatin-like lipase inhibitors	novel and useful industrial	2014)
using a taxonomy-based dereplication with a public	Streptomyces strains	
collections of strains and in vitro assays		

## Conclusion

Dereplication procedures now play a prominant role in the field of natural product research. The examination of all bibliographic references published from 1990 to 2014 which involve the concept of dereplication clearly reveals that five different dereplication strategies have been gradually developed to meet specific objectives. The common denominator between these studies is the willingness to accelerate the discovery of biologically active substances by improving characterization methods of natural resources. Most dereplication workflows have addressed this issue by using targeted or untargeted chemical profiling strategies, often in combination with biological assays and computer tools to assess the presence of active metabolites in natural samples, while others have used a taxonomic approach mainly based on genetic analyses for the classification and prioritization of samples. It is quite clear that the quality and completeness of small metabolite databases and microbial strain libraries remain the major impediments to high performance dereplication workflows. Consequently, the author can only stress the importance of finding a way to link collected spectral data between laboratories and finally homogenize these databases.

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