Novel approach for identification of biocatalysts by reverse omics techniques

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Chapter I

Summary
1. Summary

A novel screening approach for identification of biocatalysts by reverse omics techniques was developed. Basic principle is the differential analysis of metatranscriptomes obtained from cultures treated either with the substrate or the product of the enzymatic reaction of interest. All genes upregulated in the substrate-containing culture respond to the added compound and could be involved in its degradation or belong to respective transporters or resistance mechanisms.

Nitrilases, enzymes degrading often toxic nitriles to the corresponding carboxylic acids and ammonia, were chosen as biocatalysts of interest. As effect of nitriles on microbial life is barely known, their influence had to be determined before establishment of the screening approach. For that purpose, agar plates containing different concentrations of nine tested nitriles (phenylacetonitrile, succinonitrile, acetonitrile, crotononitrile, 4-hydroxybenzonitrile, acetone cyanohydrin, cyclohexanecarbonitrile, 2-phenylpropionitrile, and pyruvonitrile) were prepared. Growth of \textit{Agrobacterium tumefaciens}, \textit{Bacillus subtilis}, \textit{Corynebacterium glutamicum}, and \textit{Escherichia coli} was monitored on these plates and revealed first insights into nitrile toxicity. Subsequently, highest non-toxic concentrations were used to treat liquid cultures containing a microbial community derived from compost. Growth was monitored regularly and cultures were transferred to fresh medium every second day to avoid nutrient depletion. Acetone cyanohydrin, 2-phenylpropionitrile, and pyruvonitrile exhibited a lethal effect on the microbial community. In contrast, cultures containing succinonitrile, acetonitrile, and crotononitrile showed higher optical densities than the control, indicating a growth-supporting effect. Furthermore, community composition was determined by 16S rRNA gene analysis and metagenome sequencing, revealing specific community-shaping effects for every compound, i.e. \textit{Pseudomonas} was detected in most cultures whereas \textit{Paenibacillus} was highly abundant in cultures containing growth-suppressing nitriles. In general, Gram-positive bacteria showed higher nitrile tolerance than Gram-negative bacteria.

Growth-supporting effect of acetonitrile during nitrile toxicity analysis indicated metabolization of this compound and therefore presence of nitrile-degrading enzymes. To increase knowledge on nitrile-degrading organisms and obtain strains for establishment of the screening approach, isolation of respective organisms was performed. Finally, eight different isolates belonging to \textit{Flavobacterium}, \textit{Pseudomonas}, \textit{Rhodococcus}, and \textit{Variovorax} were identified. Nitrile-degradation by the latter three is common, but only weak degradation by \textit{Flavobacterium} has been reported before. Genome sequencing of the isolates revealed various nitrile-degrading enzymes for all strains except \textit{Flavobacterium}, indicating a novel mechanism for nitrile degradation in this genus.

Analysis of metagenomes obtained during nitrile toxicity test revealed 70 putative nitrilases in nitrile-treated cultures. Due to their origin, they were promising candidates for identification of novel nitrile-
degrading biocatalysts. For fast and simple screening of this number of enzymes, a novel high-throughput assay was developed. The new method combines real-time measurement of enzymatic activity and high sensitivity without dependency on purified proteins. Six putative enzymes exhibited nitrilase activity and subsequently, the most interesting biocatalysts was further characterized. The novel nitrilase exhibits a broad pH optimum and unusually high long-term stability. Furthermore, it is highly specific for phenylacetonitrile and belongs to the class of arylacetonitrilases, which are of industrial importance but rarely studied.

Finally, the proposed screening approach for identification of biocatalysts by reverse omics techniques was developed. For initial tests, the previously isolated acetonitrile-degrading *Rhodococcus* was used. Growth condition, cell harvesting, and isolation of DNA and RNA were optimized with this strain. Subsequently, a protocol for normalization and differential analysis of transcriptomes was established. Only one operon consisting of three genes and a nearby amidase were significantly upregulated in the nitrile-treated cultures. The operon did not encode for known nitrile-degrading proteins, but proximity to the amidase supports the respective activity. Subsequently, the established parameters were applied to a complex microbial community. Metatranscriptomes were normalized based on the abundance of single species to allow statistically valid analyses. More than 500 up- and 280 downregulated genes could be identified under nitrile-treatment, indicating complex interactions between members of the community. For one species, a highly upregulated nitrilase and amidase were found, whereas no significant upregulation of respective genes was recorded for other species. Therefore, unknown nitrile-degrading enzymes may hide behind upregulated putative proteins, demonstrating the potential of this novel metatranscriptomic screening approach.
Chapter II

General introduction
2. General introduction

During the first half of the 19th century, the era of modern chemistry began. In 1835, Jöns Jacob Berzelius first mentioned “catalysis” as a process in which a compound influences a chemical reaction without being part of the substrates or products (1). Together with observations of the catalytic reactivity of metals like platinum, palladium, and iridium (2, 3), a new research field of chemistry was created. Early industrial application of catalysts started in 1913 with the synthesis of up to 30 tons ammonia per day using an iron catalyst (4). In the following decades, numerous other catalytic processes were established, e.g. the synthesis of hydrocarbons using carbon monoxide and hydrogen, the synthesis of methanol, the polymerization of ethylene to polyethylene, and cracking of petroleum using zeolite as catalyst (5).

Even though modern civilization would not be possible without chemical industry, large scale production results in a severe environmental footprint. Many chemical processes need strong acids or bases, (heavy) metal catalysts, high reaction temperature, and produce large amounts of inorganic waste. In addition, the processes can yield unwanted isomers (6–8). For example, the production of 1 ton ammonia leads to the release of 1.5 tons carbon dioxide. In addition, global ammonia production is responsible for 2% of the total industrial energy consumption (9). Subsequently, ammonia can be used for production of compounds like acrylonitrile. More than 7 million tons of this important monomer are produced per year (10), but synthesis depends on high temperature and pressure. In addition, several purification steps for removal of contaminants like hydrogen cyanide or acetone are necessary to yield pure acrylonitrile (11).

With a growing population and increasing standards in developing countries, limited resources and environmental problems like climate change become more challenging. Therefore, chemical industry must advance and develop new environmentally friendly processes to fulfill the growing demand for bulk and fine chemicals.

2.1. Green chemistry

During the 1980s, waste production and use of toxic or hazardous materials in industry became a growing concern (12), finally leading to the U.S. Pollution Prevention Act of 1990 (13). In contrast to previous governmental guidelines it focused on pollution prevention rather than waste treatment, marking the beginning of the green chemistry era. Subsequently, twelve principles of green chemistry were developed (14):

1) Waste should be prevented instead of being treated

2) Incorporation of all involved materials into the final product should be maximized
3) Use or generation of toxic substances during production should be minimized
4) Toxicity of final products should be reduced
5) Use of auxiliary substances like solvents should be avoided
6) Energy requirements of chemical processes should be minimized
7) Renewable feedstock or raw material should be used when practical
8) Unnecessary derivatization should be minimized
9) Catalytic reagents should be preferred over stoichiometric reagents
10) Products should break down into innocuous products at the end of their use
11) Real-time analyses should be further developed to prevent pollutant formation
12) Inherently safer chemicals should be used for accident prevention

These principles are nowadays often used as guideline for the development of new production lines. The most promising approach for a sustainable green chemistry is the transition to biobased processes, as demonstrated by novel processes for production of acrylonitrile. Traditional synthesis is based on ammonia and propene, a byproduct of oil refining, and therefore not sustainable. In contrast, newly developed synthesis routes use renewable substrates like glycerol or glutamic acid (15). Furthermore, they can achieve higher yields and avoid by-products like hydrogen cyanide (10). Nevertheless, not all aims of green chemistry can be fulfilled by change of substrates, as these novel processes still need unfavorable conditions and metal catalysts. Consequently, enzymes as biocatalysts are of increasing interest as they can help to meet 10 of the 12 criteria of green chemistry (16).

2.2. Biocatalysis and bioremediation

Although the principles of catalysis were almost unknown, the potential of enzymes was already discovered in the 19th century, when Louis Pasteur demonstrated the conversion of L-(-)-tartaric acid to D-(-)-tartaric acid using *Penicillium glaucum* (17). In 1874, a Danish company (Christian Hansen’s Laboratory) started selling enzyme preparations for the production of cheese (18). In 1897, Eduard Buchner showed that living organisms are unnecessary for enzymatic activity by using cell-free yeast extract for fermentation of sugar, further increasing the interest on biocatalysts (19). Enzymes received attention by biologists, but chemists did not recognize their potential as (bio)catalysts (20). Even the immobilization of enzymes and a similar handling to classic chemical catalysts did not change this view (21). One of the main reasons for this difference may have come from limitations during enzyme isolation. As natural producers were often difficult to grow, only small isolation volumes and therefore irrelevant quantities for industrial purposes were obtained. This problem was solved with development of recombinant DNA techniques in the 1970s and PCR in 1983. Transfer of interesting genes into heterologous easy-to-handle host organisms like *Escherichia coli* or
Bacillus subtilis suddenly allowed the efficient production of enzymes (22). In addition, development of directed mutagenesis and directed evolution allowed enhancements of substrate range or protein stability, finally leading to industrially relevant enzymes (20, 23, 24).

Modern biocatalysts show high selectivity, work under mild conditions, and can be used without the production of critical waste (25). Especially the pharmaceutical industry has interest in biocatalysts as many conventional drug synthesis processes produce large amounts of waste (26). The early use of a lyase for the production of L-3,4-Dihydroxyphenylalanin, an important drug for the treatment of Parkinson’s disease, underlines this interest (27). Nowadays, application of biocatalysts is widespread. The food industry uses enzymes for bread and cheese production (28), whereas bulk production of bioethanol from starch (42 million m³ per year) demonstrates the full potential of enzymatic processes (21). The huge industrial interest reflects also on market data: In 2018 the global enzyme market had a value of US $5.5 billion, but is expected to reach US $7.0 billion in 2023 (29).

With rising awareness of environmental pollution in the public, waste prevention and remediation became growing industrial sectors. Bioremediation aims to remove pollution appearing during production processes or remediate already contaminated environments by using (genetically modified) microorganisms. Examples are the biosorption of metals like nickel, lead, or cadmium from soil or wastewater (30, 31) or the bioremediation of radionucleotide-contaminated sites (32, 33). Furthermore, microorganisms can be used for removal of mutagenic or carcinogenic dyes from textile industry wastewater (34, 35). In rare cases, it can even be the only option for environmental decontamination. During an earthquake in India in 2001, damaged acrylonitrile tanks led to the release of the chemical and contamination of surrounding soil. Eight months after leakage, no reduction of nitrile concentration could be measured, demonstrating its persistence. Subsequent bioaugmentation and nutrient-supplementation helped to establish microbial soil communities with nitrile-degrading properties, finally leading to successful decontamination of the environment (36, 37).

2.3. Natural nitriles and their toxicity

Nitriles are toxic compounds with \(-\text{C}=\text{N}\) as functional group. Most nitriles are synthesized from amino acids (tyrosine, phenylalanine, valine, isoleucine, and leucine) which are converted to aldoximes and subsequently transformed to nitriles or cyanogenic glycosides (Figure 1) (38–40). Nitriles are widespread in nature and have been identified in form of hydrogen cyanide in insects (41, 42), as aeroplysinin in sponges (43), or as antibiotics like borrelidin or toyocamycin in bacteria (44, 45). Furthermore, several nitrile-containing compounds have been reported for fungi (46–48). The most prominent group of nitriles are cyanogenic glycosides, which can be found in over 3,000 plant species belonging to more than 130 families (49).
Most of the naturally occurring nitriles and cyanides seem to be involved in defense mechanisms. Toyocamycin as nucleoside antibiotic interferes with cell wall synthesis (45) and borrelidin inhibits the activity of threonyl-tRNA synthetases (50). For aeroplysinin, several protective functions like antibiotic, anti-inflammatory, or anti-angiogenic effects have been described (51). Main purpose of cyanogenic glycosides seems to be defense against predators by degradation to hydrogen cyanide after cell disruption or attack (52, 53), but use as nitrogen storage has also been described (54).

![Diagram of nitrile synthesis](image)

**Figure 1: Nitrile synthesis in nature.** Adapted from Prasad et al., 2010 (38).

Although many nitriles have toxic effects, toxicity does not derive from the nitrile structure itself. The proposed mechanism in eukaryotes assumes nitrile degradation by cytochrome P450 to an unstable cyanohydrin. Spontaneous decomposition of this compound leads to the release of highly toxic cyanide, which can inhibit cytochrome c oxidases (55–59). As this enzyme is the terminal electron acceptor of the respiratory chain, inhibition leads to asphyxiation and can be lethal. Even though respective studies for bacteria are missing, a similar mode of action can be assumed as cytochrome P450 and cytochrome c oxidase are present in many bacteria (60, 61).

Besides cyanide release, other mechanisms for toxicity are also known. For example, main toxicity of acrylonitrile arises from interaction with thiol groups, affecting central and peripheral nervous systems (62). As thiol groups are also important for activity of numerous bacterial enzymes, similar toxicity mechanisms may apply. Other mechanisms may also account for toxicity of chloroacetonitrile, isovaleronitrile, caprylonitrile, and benzonitrile, as no linkage between nitrile concentration and intracellular cyanide presence could be found (55). In addition, chemical structure seems to have an influence on toxicity. An increased chain length appears to reduce toxicity and dinitriles are slightly
less toxic than mononitriles. Nevertheless, toxicity values observed for the first nitrile of a homologous series cannot be extrapolated to other members of such a group (63).

2.4. Nitrile-degrading enzymes

As nitriles and cyanides are toxic to living organisms, different enzymatic detoxification strategies exist (Figure 2). For degradation of cyanide, mainly two pathways are used. The first involves cyanide hydratases (EC 4.2.1.66) catalyzing the degradation of cyanide to formamide. They are most likely exclusively fungal enzymes consisting of several subunits and can reach up to 3,000 kDa (64–66). The second pathway uses cyanide dihydratases that degrade cyanide to formic acid and ammonia. So far, this reaction was only detected in a few bacterial genera (67). Like cyanide hydratases, cyanide dihydratases do not require metal cofactors and are therefore interesting candidates for bioremediation of cyanide-contaminated sites (68). In addition to these two options, other pathways such as direct degradation to carbon dioxide and ammonia by cyanide dioxygenases or to methane and ammonia by nitrogenases are known (69).

Figure 2: Enzymatic nitrile degradation. Adapted from Prasad et al., 2010 (38).
Degradation of nitriles is mainly performed by two pathways. The first involves nitrilases (EC 3.5.5.1), which degrade nitriles directly to carboxylic acid and ammonia. These enzymes are known in archaea, bacteria, fungi, plants, and animals (70) and belong to the carbon-nitrogen hydrolase superfamily. Nitrilases are classified according to their preferred substrates into aliphatic, aromatic, and arylacetonitrilases (71). They consist of 6-26 subunits with an α-β-β-α fold and an average mass of 40 kDa (72–74). Important for enzymatic activity is a metal-independent conserved catalytic triad which is present in all known members of the nitrilase family and consists of glutamic acid, lysine, and cysteine (72, 75–78). Majority of these enzymes have a temperature optimum of activity between 30 °C and 55 °C and show catalytic activity under neutral or slightly alkaline conditions (74). Their expression is normally induced by their substrates or products, but a few cases of constitutive nitrilase expression have also been described (79–81).

The second pathway for nitrile degradation consists of two enzymes, nitrile hydratases (NHases; EC 4.2.1.84) and amidases (EC 3.5.1.4). The former degrade nitriles to amides and the latter subsequently the amides to carboxylic acid and ammonia (82). In contrast to nitrilases, NHases belong to the carbon-oxygen lyase superfamily and are found in several bacterial genera and few eukaryotes (38, 83). They consist of an α- and β-subunit with average molecular masses of around 25 kDa (38, 84, 85). Based on the metal cofactor bound to their α-subunit, these enzymes are often divided into cobalt or ferric NHases (86). Besides the cofactor, conserved amino acids seem to be important for catalytic activity. For cobalt NHases, respective threonine and tyrosine residues have been identified, whereas specific serine and threonine residues are conserved in ferric NHases (87, 88). In addition, it has been shown that at least for a NHase from Streptomyces rimosus the α-subunit is sufficient for enzymatic activity (89). In contrast to nitrilases, NHases are often thermolabile and exhibit highest activity between 30 °C and 40 °C, but share the same optimal pH range at around pH 7 (38).

Amidases, the second enzyme of this bi-enzymatic pathways, are similar to nitrilases. They also belong to the carbon-nitrogen hydrolase superfamily and contain the same conserved catalytic triad (90). These ubiquitous enzymes have an average molecular mass of around 40 kDa and can consist of monomers, dimers, tetramers, hexamers, or octamers (91–94). Except for amidases involved in cell separation using zinc, cofactors are not required for enzymatic activity (95). Most of the characterized amidases show optimum activity at neutral or slightly alkaline pH values and temperatures between 40 and 55 °C (91, 96–99).
2.5. Nitriles and nitrile-degrading enzymes in industry

Nitriles as well as nitrile-derived amides and carboxylic acids are of great importance for industry. The first amidase was described in 1949 (100), followed by nitrilases in 1964 (101), and NHases in 1980 (102). Increasing numbers of publications for all these enzymes demonstrate the intensified research activity in this area (Figure 3). Nevertheless, literature on nitrile-degrading enzymes is scarce when compared to other enzymes like phosphatases, esterases, or lipases.

![Figure 3: Total number of publications targeting nitrile-degrading enzymes. Data were obtained by search for “amidase”, “nitrile hydratase”, and “nitrilase” in the Web of Science. Date of search: 12th November 2018.](image)

Nitriles and nitrile-derived products can be found in different fields of industry. In pharmacy, more than 30 drugs containing nitrile groups are used nowadays and several more are in clinical studies. Their spectrum reaches from antidiabetics (vildagliptin) and anticancer drugs (anastrozole) over cardiotonic agents (olprinone) to antidepressants (escitalopram) (103). Probably the most important application in pharmacy is the utilization of a penicillin amidase to produce 6-aminopenicillic acid from penicillin G, allowing the cost-efficient production of penicillin derivates (104, 105). In addition, a nitrilase catalyzing the hydrolysis of the industrially produced 2-(4’-isobutylphenyl)propionitrile to ibuprofen is known and leads to high yields of the active isomer (106).

Besides synthesis of small amounts of expensive chemicals, nitriles and nitrile-degrading enzymes are also used for production of bulk chemicals. Companies like BASF AG or Mitsubishi Rayon use NHases for production of more than 600,000 tons acrylamide per year (107, 108). Another example for industrially used NHases is the production of 11,500 tons nicotinamide, a form of vitamin B3, by Lonza in 2006 (109). Furthermore, hundreds of tons of mandelic acid and nicotinic acid are produced per year.
using nitrilases (74, 110). Especially the synthesis of nicotinic acid is a good example how biocatalysts can help to fulfill the principles of green chemistry. The most common manufacturing process is based on liquid-phase oxidation of 2-methyl-5-ethylpyridine at high temperatures and high pressure (111). In addition, it depends on the use of toxic, flammable, and hazardous nitric acid, which is difficult to handle and a threat to the environment (112). With transition to nitrilase-based production processes, these unfavorable conditions are omitted, leading to a safer and greener chemical industry.

Nitriles are not only industrially important substrates or products, but also used in agriculture. Prominent examples are the herbicides bromoxynil and dichlobenil (113, 114), the latter also available for private households. Transgenic oxynil-herbicide resistant crops have been developed to deal with toxic effects (115) but accumulation in the environment could cause problems. In addition, other industrially used nitriles released into (waste)water and soil have deleterious effects on the local biota. Consequently, bioremediation involving nitrile-degrading enzymes is of increasing scientific and economic interest. Classic treatment of nitrile-contaminated industrial waste is performed with chemicals like hydrogen peroxide or by alkaline chlorination, carrying the risk of subsequent contamination with toxic chemicals (116). Therefore, several remediation processes based on enzymatic nitrile degradation have been developed during the last decades. Most of them focus on immobilization of nitrile-degrading bacteria with alginate to simplify operation of decontamination systems (117, 118). Another approach uses bacterial consortia consisting of biofilm-forming and nitrile-degrading organisms for bioremediation of acetonitrile-containing waste water (119). In addition, a recombinant biofilm-forming Bacillus containing a Rhodococcus nitrilase was created (120). Due to its biofilm, this strain exhibits high resistance to nitrile toxicity and is therefore a promising organism for industrial bioremediation.

### 2.6. Enzyme screening methods

Discovery of novel enzymes is important, especially as not every limitation for industrial application can be solved by protein engineering. Probably the most traditional screening methods are culture-based techniques. For that purpose, a microbial inoculum is grown under presence of a chemical of interest used as e.g. sole carbon source. Only microorganisms with desired enzymatic can grow by metabolization of the added compound and are further analyzed. Similarly, agar plates containing indicator dyes can be used, showing color changes or halos around microorganisms with desired activity (121, 122). Despite its simplicity, this technique is highly limited as only 1 % of all microorganisms can be grown under standard laboratory conditions. Therefore, most metabolic potential cannot be harvested by this approach (123).
As consequence, two screening approaches have been developed to explore the genetic diversity of non-culturable microorganisms. The first method uses gene libraries from metagenomic DNA and activity-based screens. By combining expression of metagenomic genes in hosts like *Escherichia coli* with simple activity assays like the previously described plate method, novel enzymes can be identified (124–127). The drawback of this system is its high effort and problems with the heterologous expression, as often less than 50% of the investigated genes are expressible in *E. coli* (128).

The second method uses comparative genomics to mine (meta)genomes for new biocatalysts. For that purpose, (public) sequence data are searched for homologues of an enzyme of interest, often leading to the prediction of several new proteins. Subsequently, these candidates are subcloned into *E. coli* or other hosts and tested for activity. Sequence-driven discovery of many novel enzymes such as methyl halide transferases, mandelate oxidases, rare arylmalonate decarboxylases, and nitrilases demonstrates the potential of this approach (129–132). Nevertheless, identified putative proteins do not necessarily exhibit the targeted function. Furthermore, entirely novel enzymes cannot be identified as only homologues to already known proteins are detected (133).

Besides (meta)genome-based screening approaches, transcriptome-based screenings have been developed recently. By comparison of different growth conditions, differential transcriptomics can help to identify novel enzymes or metabolic pathways. This approach was already used for determination of genes involved in heat-shock response in *Brevibacillus borstelensis*, adaption of *E. coli* to different toxins, or identification of biocatalysts for synthesis of chiral alcohols in *Pseudomonas monteilii* (134–136). Major benefit of this approach is the simultaneous detection of several interesting genes at once, also allowing the identification of novel enzymes and entire pathways. Nevertheless, differential transcriptomics for identification of biocatalysts have only been applied to single organisms. Therefore, neither complex interactions between members of microbial communities nor organisms difficult to isolate can be analyzed by this approach.

2.7. **Aim of the thesis**

Aim of this thesis is the establishment of a novel approach for identification of biocatalysts by reverse omics techniques. This approach is based on differential analysis of metatranscriptomic datasets without the necessity of designing a suitable screen for the targeted biocatalyst. For this purpose, microbial communities are incubated in the presence and absence of the target compound. Subsequently, mRNA of both approaches is isolated and sequenced. Ideally for quantification and later identification of full-length genes, the RNA datasets are mapped on a metagenome backbone determined from the same sample. By differential analyses of the metatranscriptomic dataset, genes specifically expressed under the presence of the added compound are identified.
Nitrilases were chosen as genes of interest. Therefore, growth media for the proposed metatranscriptomic screening approach contain either a nitrile or its degradation product in form of the corresponding carboxylic acid and ammonia. Upregulated genes identified during differential analysis of respective metatranscriptomes should belong to nitrilases, nitrile hydratases, amidases, or unknown resistance mechanisms. Before testing the novel screening approach, toxicity of different nitriles and corresponding carboxylic acids must be determined to avoid unintended lethal effects during screening. With the respective values in hand, the novel screening approach will be tested. A known nitrile-degrading organism will be used to establish growth conditions, sample preparation, and bioinformatic analysis. Subsequently, the optimized parameters will be applied to a complex microbial community, leading to the identification of novel biocatalysts.
Chapter III

Results
3.1. Impact of nitriles on bacterial communities

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\textit{Supplementary data can be found on the enclosed DVD}

Author contributions

Performed the experiments: \textbf{RE, TZ}

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Conceived the study: \textbf{RH, RD}
**ABSTRACT**

Nitriles are organic molecules with a nitrile group (−C≡N) as functional group and often toxic for living organisms. Detoxification can occur via nitrilases that degrade nitriles directly to carboxylic acids and ammonia, or with nitrile hydratases and amidases that convert nitriles to amides and subsequently to carboxylic acids and ammonia. Despite the knowledge of enzymatic degradation pathways, the influence of these compounds on the composition of bacterial communities has not been studied. Initially, the tolerances of four phylogenetically different bacterial strains without known nitrile detoxification systems (Agrobacterium tumefaciens, Bacillus subtilis, Corynebacterium glutamicum, and Escherichia coli) to the toxic effects of nine different nitriles and the corresponding carboxylic acids were determined. Based on these results, the effect of nitriles on diversity and composition of environmentally-derived bacterial communities was monitored over time by 16S rRNA gene amplicon-based and metagenome analyses. Three nitriles (acetone cyanohydrin, 2-phenylpropionitrile, and pyruvonitrile) exhibited a lethal, three (phenylacetonitrile, 4-hydroxybenzonitrile, and cyclohexanecarbonitrile) a growth-suppressing and three (succinonitrile, acetonitrile, and crotononitrile) a growth-promoting effect on the studied communities. Furthermore, each tested nitrile exhibited a specific community-shaping effect, e.g. communities showing growth-suppression exhibited high relative abundance of Paenibacillus. In general, analysis of all data indicated a higher resistance of Gram-positive than Gram-negative bacterial community members and test organisms to growth-suppressing nitriles. Finally, more than 70 putative nitrilase-encoding and more than 20 potential nitrile hydratase-encoding genes were identified during analysis of metagenomes derived from nitrile-enrichments, underlining the high yet often unexplored abundance of nitrile-degrading enzymes.

**IMPORTANCE**

In this study, we present a systematic analysis of the influence of different nitriles on a complex microbial community. Significant changes of bacterial community diversity and composition depending on the added nitrile occurred. Furthermore, numerous putative nitrile-decomposing enzymes were identified from metagenomes. Thus, the here presented results reveal the potential impact of nitriles and their degradation products on microbial communities and approaches for its determination.
INTRODUCTION

Nitriles are structurally diverse organic molecules with a nitrile group (−C≡N) as functional group. Naturally occurring nitriles are found in bacteria, fungi, plants, and animals as well as in marine and terrestrial habitats (1–4). The most widespread nitriles are cyanogenic glycosides, which often are present in plants (5) and other organisms (6). They can serve as storage form of reduced nitrogen (7) or defense compound when degraded to highly toxic hydrogen cyanide (4).

Toxicity of most nitriles does not originate from the nitrile itself, but from cyanide release due to degradation by e.g. cytochrome P450 in eukaryotes (8–10). Subsequently, cyanide inhibits cytochrome c oxidases and therefore the terminal electron acceptor of the respiratory electron transport chain (11). As cytochrome P450 and cytochrome c oxidase are also present in many bacteria (12, 13), a similar mode of action as in eukaryotic organisms is indicated.

Detoxification and degradation are facilitated by two known enzymatic routes. The first involves nitrilases (EC 3.5.5.1), which belong to the carbon-nitrogen hydrolase superfamily and harbor a conserved catalytic triad consisting of glutamic acid, lysine, and cysteine (14, 15). These enzymes degrade nitriles directly to carboxylic acids and ammonia. In the second route, nitriles are degraded to amides and subsequently to carboxylic acids and ammonia. The hydration of nitriles to amides is catalyzed by nitrile hydratases (NHases; EC 4.2.1.84), which consist of an α- and β-subunit and are classified based on the used cofactor as cobalt or ferric NHases. The subsequent hydrolysis of amides to carboxylic acids and ammonia is performed by amidases (EC 3.5.1.4) (16).

Nitriles are also important industrial precursors for the production of bulk chemicals like acrylamide (17) or part of various pharmaceuticals such as the anti-cancer drug letrozole or etravirine for HIV treatment (18). Non-intended release of man-made nitriles can lead to environmental accumulation and impact natural communities. In addition, some nitriles are used as herbicides, e.g. dichlobenil (19) or bromoxynil (20). The effect of these synthetic compounds on bacterial communities, the key players in most nutrient cycles (21–23), is almost unknown. While toxicity values and effects of many nitriles on mammals have been reported, comparable values are almost unavailable for microorganisms (8, 11, 24, 25).

The aim of this work was to investigate the effect of nine nitriles and their corresponding degradation products on bacterial communities. Initially, the toxicity of these nitriles and their corresponding carboxylic acids were explored using four bacterial test strains containing no known nitrile detoxification system. Based on these results a community growth experiment was performed and revealed the specific impact of each nitrile on a bacterial community with respect to composition and diversity. Additionally, metagenome sequences provided insight into potential mechanisms of nitrile stress management, as quantity and distribution of nitrile-degrading enzymes were analyzed.
RESULTS

Limits for bacterial life. To establish stable nitrile-containing mixed cultures it is necessary to determine the toxic effect of these compounds. Therefore, we analyzed the influence of a set of nine nitriles on the four model organisms *Agrobacterium tumefaciens*, *Bacillus subtilis*, *Corynebacterium glutamicum*, and *Escherichia coli* by using agar plates containing a rich complex or a minimal medium with different nitrile concentrations. All four tested strains do not contain any known nitrile-degradation pathway. As carboxylic acids, the products of a potential nitrile detoxification reaction, also affect microbial growth, their influence was tested in addition.

Growth was monitored for four days on solid agar plates supplemented with a nitrile or the corresponding carboxylic acid and examined every 24 hours (Data Set S1). In general, the Gram-positive strains (*B. subtilis* and *C. glutamicum*) showed better growth at higher nitrile and carboxylic acid concentrations than the Gram-negative strains (*A. tumefaciens* and *E. coli*). Some tested strains revealed different behavior with respect to the nitriles and the corresponding carboxylic acids. For example, 4-hydroxybenzonitrile suppressed the growth of *A. tumefaciens* and *E. coli* even at 5 mM concentration but all organisms except *A. tumefaciens* were able to grow at 25 mM of the corresponding carboxylic acid (Table 1). The higher susceptibility of the tested Gram-negative strains was also observed for pyruvonitrile, as growth was inhibited at 15, 20, and 25 mM concentration, whereas Gram-positive strains grew at 15 mM. In addition, cyclohexanecarbonitrile had no influence on the growth of all tested organisms, but cyclohexanecarboxylic acid inhibited growth at 10 mM.

The results provided a first insight into the inhibitory potential of the nitriles and carboxylic acids with respect to bacterial growth. The highest non-inhibitory concentration considering all four test strains was chosen for further experiments. In cases where the highest concentration of a nitrile and the corresponding carboxylic acid differed, the lower concentration was chosen for both substances. For substances inhibiting growth even at lowest tested concentration, 5 mM were used subsequently.
### Table 1: Bacterial growth with nitriles and corresponding carboxylic acids after four days of incubation

Effect of nine different nitriles and corresponding carboxylic acids at 5, 10, 15, 20, and 25 mM in LB and M9-HEPES on the growth of *A. tumefaciens*, *B. subtilis*, *C. glutamicum*, and *E. coli* after four days of incubation. 0, no growth; 1, *A. tumefaciens* growing; 2, *B. subtilis* growing; 3, *C. glutamicum* growing; 4, *E. coli* growing. Underlined numbers represent the concentration chosen for growth of environmental sample.

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<th>LB 10 mM</th>
<th>LB 15 mM</th>
<th>LB 20 mM</th>
<th>LB 25 mM</th>
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<th>M9-HEPES 10 mM</th>
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Nitrile and carboxylic acid impact on bacterial growth. To determine bacterial community alterations under the influence of nitriles and their corresponding carboxylic acids a growth experiment in liquid minimum medium was performed. A compost sample from the Experimental Botanical Garden Göttingen (Germany) served as inoculum.

A general reduction of the final OD with each cultivation passage was observed for the nitrile-based and the carboxylic acid-based cultures (Figure 1). At the last measuring point of the carboxylic acid-supplemented cultures, seven of nine conditions revealed higher optical densities than the control, thereby indicating beneficial effects of the supplemented carboxylic acids. In case of the nitrile-supplemented cultures only three nitriles (succinonitrile, acetonitrile, and crotononitrile) exhibited a growth-supporting effect in comparison to the control without nitriles. The beneficial impact of acetonitrile was already obvious from the second passage on. Addition of cyclohexanecarbonitrile did not affect growth, whereas a negative effect on microbial growth was observed for phenylacetanitride and 4-hydroxybenzonitrile. In contrast, cultures supplemented with acetone cyanohydrin, 2-phenylpropionitrile, or pyruvonitrile showed no growth at all after the first passage.

Figure 1: Growth of a microbial community in M9-HEPES treated with (A) nitriles or (B) the corresponding carboxylic acids.
**Bacterial community composition and diversity.** The bacterial community composition of the control cultures without addition of nitriles or the corresponding carboxylic acids was determined via amplicon-based 16S rRNA gene analysis (Fig. 2 and Data Set S2). Analysis of Simpson indices of diversity (SID) at genus level revealed reduction of diversity from inoculum (SID 0.98) to the first sampling of the control after 68 h of incubation (SID 0.58). Diversity increased again after 118 h of incubation (SID 0.78) and stabilized in the later taken control samples (166 and 214 h; SID 0.91 in both cases). Besides inoculum and control cultures, only communities showing growth after 214 h with the nitrile or the corresponding carboxylic acid were analyzed by 16S rRNA gene analysis (Figure 2). Cultures supplemented with acetone cyanohydrin, 2-phenylpropionitrile, and pyruvonitrile were excluded from this analysis due to absence of viability after first passage. Consequently, the respective carboxylic acid-supplemented cultures were also removed, as in these cases comparison between a nitrile and its carboxylic acid was not feasible.

![Figure 2: Abundance of bacterial genera based on 16S rRNA gene analysis in cultures containing different nitriles or carboxylic acids after 214 h of incubation.](image)

Nitrile-supplemented cultures indicated a general formation of two distinct bacterial community types. Cultures containing succinonitrile, acetonitrile, or crotononitrile showed a high diversity (SID 0.88, 0.91 and 0.86, respectively) when compared to cultures with phenylacetonitrile, 4-hydroxybenzonitrile or cyclohexanecarbonitrile (SID 0.52, 0.14 and 0.65, respectively). Cultures supplemented with the corresponding carboxylic acids exhibited a similar pattern. Communities with acetic acid (SID 0.77), crotonic acid (SID 0.73), 4-hydroxybenzoic acid (SID 0.82), and
cyclohexanecarboxylic acid (SID 0.83) exhibited a higher diversity compared to cultures supplemented with phenylacetic acid (SID 0.61) and succinic acid (SID 0.56). In addition, some carboxylic acid-containing cultures showed strong variations in diversity in relation to the corresponding nitrile-containing cultures. In case of succinonitrile and succinic acid, diversity decreased from SID 0.88 to 0.56. A similar trend was observed for acetonitrile (SID 0.91) and acetic acid (SID 0.77). In contrast, a highly increased diversity could be detected in 4-hydroxybenzoic acid-containing cultures (SID 0.82) compared to the corresponding nitrile (SID 0.14). These differences in diversity indicate a specific effect of the nitrile group in comparison to the carboxylic acid group.

Nitrile-affected cultures showed distinct communities when compared with the carboxylic acid counterparts. *Pseudomonas* was present under most conditions, except in cultures containing phenylacetonitrile or 4-hydroxybenzonitrile. A similar behavior was recorded for *Aeromonas*. Interestingly, *Paenibacillus* is the dominant genus in phenylacetonitrile- or 4-hydroxybenzonitrile-containing cultures with relative abundances of 57% and 92%, respectively. Members of this genus are also present in all other cultures but in lower relative abundances. In addition, *Brevundimonas* was present (>1% relative abundance) in three of the six nitrile-treated cultures, but only in one containing a carboxylic acid. Furthermore, *Acinetobacter* and *Aeromonas* showed an on average higher relative abundance under acetic acid treatment. In contrast, *Pedobacter* was only identified in the controls (>1% relative abundance), while *Aerophila*, *Chryseobacterium*, and *Glutamicibacter* were only present after treatment with phenylacetonitrile, succinonitrile or 4-hydroxybenzonitrile, respectively. Finally, *Serratia* showed high relative abundance in cultures containing phenylacetonitrile (40 %) or crotonic acid (42 %).

The combination of the 16S rRNA gene analysis with the corresponding growth curves revealed further linkages between the density of a culture and its diversity. Cultures with phenylacetonitrile and 4-hydroxybenzonitrile did not only reveal reduced optical density, but a reduced diversity when compared to the cultures containing the corresponding carboxylic acids. In contrast, succinic acid-containing cultures showed low diversity, but exhibited higher density than the control. Finally, several cultures (succinonitrile, acetonitrile, 4-hydroxybenzoic acid, and cyclohexanecarboxylic acid) showed higher density and higher diversity than the control, indicating a growth-benefiting effect of the added chemicals.
How to handle nitriles. To assess in which way the different bacterial communities manage the supplemented nitriles, metagenomes of all cultures incubated for 214 h and showing growth were sequenced. Subsequently, the metagenomes were assembled, annotated, and analyzed. The samples were checked for presence of genes encoding nitrilases and nitrile hydratase subunits. Finally, the taxonomic assignment of the contigs harboring the targeted genes was used to infer the phylogenetic origin of the deduced enzymes (Data Set S3).

In general, the metagenomes showed a similar taxonomic composition as previously determined via 16S rRNA gene analysis. Other domains were barely present, as only three respective microorganisms were detectable (Figure 3A). The first is the nematode *Parastrongyloides*, whose DNA was detected in the control culture after 166 h of incubation (2 %) as well as in cultures with succinonitrile (4 %), acetonitrile (4 %), and crotononitrile (8 %). The second is a *Spounalikevirus* with a relative abundance of 8 % in the culture containing 4-hydroxybenzonitrile. Third, a green alga belonging to the genus *Pyramimonas* was detected in cyclohexanecarbonitrile-containing cultures with 2 % relative abundance.

Analysis of nitrile-degrading enzymes encoded by the metagenomes revealed a broad distribution. Especially members of the genus *Pseudomonas* harbor several potential genes encoding nitrilases and NHases in nitrile- and carboxylic acid-supplemented cultures (Figure 3A). Furthermore, the untreated control cultures showed high abundance of nitrile-degrading enzymes when compared to other treatments (Figure 3B).

Regarding a potential impact of a nitrile on a bacterial community, several effects were observed. For example, the abundance of *Pseudomonas* increased in the cyclohexanecarbonitrile-containing culture compared to its carboxylic acid counterpart, while the diversity of putative nitrilases was reduced by 50% (Fig. 3A). At the same time, abundance of nitrilase genes increased almost fourfold, indicating nitrile-induced enrichment of nitrile-converting genes and correspondingly enrichment of nitrile-degrading capacity of the community (Figure 3B).
Figure 3: Putative nitrilases and nitrile hydratases found in metagenomes of enrichment cultures. A, Distribution of enzymes. Numbers in white box represent number of nitrilases and numbers in black box represent number of nitrile hydratase subunits. Others, genera with an abundance of >1%. B, Gene abundance based on coverage.
Figure 4: Phylogenetic tree containing all putative metagenome-derived nitrilases. 500 bootstrap replicates were calculated and tree nodes were condensed at values below 50%. ALIPH, aliphatic nitrilase; ARO, aromatic nitrilase; ARY, arylacetoniitrilase; ALA(CN), β-cyano-L-alanine-nitrilase; CN, cyanide dihydratase. Green, putative cluster of cyanide dihydratases; blue, putative cluster of aromatic nitrilases; red, putative cluster of arylacetonitrilases; orange, putative cluster of β-cyano-L-alanine-nitrilases.

A phylogenetic tree containing all identified and reference nitrilases (Figure 4) revealed no treatment-specific clusters. Approximately 25% of all putative enzymes group together with β-cyano-L-alanine-degrading nitrilases. In addition, small clusters of arylacetonitrilases and cyanide dihydratases were found, containing 21 or 8 of the putative nitrilases, respectively. For the remaining 156 putative nitrilases, specific clustering was not recorded.
**DISCUSSION**

Studies on the susceptibility of prokaryotes to several bioactive compounds such as antibiotics (26), cytostatics (27), fungicides (28), insecticides (29), and nematicides (30) are performed frequently. To our knowledge literature addressing bacterial sensitivity towards nitriles and their degradation products has not been published. Toxicity data, determined for *A. tumefaciens*, *B. subtilis*, *C. glutamicum* and *E. coli*, indicate several differences in their susceptibility. Both Gram-positive strains revealed increased tolerance to possible growth-inhibiting effects of nitriles compared to the Gram-negative strains, whereby *A. tumefaciens* was most susceptible.

Among the tested compounds, 4-hydroxybenzonitrile was the most toxic. As the closely related compounds 3,5-diodo-4-hydroxybenzonitrile (ioxynil) and 3,5-dibromo-4-hydroxybenzonitrile (bromoxynil) are common herbicides (31, 32), a certain toxicity to microorganisms could be inferred. Especially bromoxynil has a documented negative impact on the soil microbial biomass (33). The second most toxic chemical was pyruvonitrile, which has structural similarities to the key metabolite pyruvate. This similarity leads to interference with central metabolism and deleterious effects. The toxicity of the carboxylic acids was generally reduced compared to the corresponding nitriles, except for cyclohexanecarboxylic acid. Its toxicity was significantly higher for the tested prokaryotes than that of its nitrile counterpart. The respective mechanism remains unknown but underlines the importance to investigate also potential degradation products of a nitrile.

The toxicity assay served mainly as an initial test to define working concentrations for the investigation of bacterial communities. Optical density data, representative for the viability of a community, underlined the value of the liquid culture setup. In case of the tested nitriles, only succinonitrile, acetonitrile and crotononitrile (Fig. 1A) exhibited a positive effect on microbial growth. This indicates that the predominant bacterial community members of these cultures possess the ability to degrade and utilize these nitriles. For acetonitrile this effect was already experimentally confirmed through the successful isolation of nitrile-degrading strains from similar cultures (34). All other nitrile-containing cultures revealed reduced density indicating reduced degradation ability or tolerance levels with respect to the tested compounds.

For the differences between the solid and liquid cultures with the substrates acetone cyanohydrin, 2-phenylpropionitrile, and pyruvonitrile, two explanations could be assumed. (1) In case of cyanohydrins, spontaneous decomposition has been described under different conditions (35, 36), and probably similar mechanisms also apply for other nitriles. Consequently, different stabilities of the nitriles under both tested conditions could have had influence on their effect. (2) The compost sample did not contain any acetone cyanohydrin, 2-phenylpropionitrile or pyruvonitrile resistant or degrading microorganisms. However, 16S rRNA gene analysis of the inoculum revealed the presence of
**A. tumefaciens, B. subtilis, C. glutamicum, and E. coli**, species successfully grown during the initial plate test and thus making the first assumption more likely. Furthermore, degrading organisms or at least nitrilases for acetone cyanohydrin and 2-phenylpropionitrile have already been identified (37–44), indicating the existing potential to handle these chemicals.

Monitoring diversity of the complex communities via 16S rRNA gene analysis revealed a reduction of diversity in the liquid cultures compared to the inoculum. As often more than 99% of all environmental microorganisms cannot be cultivated under laboratory conditions (45–47), a decline was expected. Generally, the 16S rRNA gene analysis of the control samples confirmed the experimental setup as appropriate and sufficient to establish a stable community. Re-diversification of the controls over the course of the experiment is explained by the prolonged incubation time. Extended incubation is a well-known factor for increasing the number of isolates on solid media (46). Apparently, regular transfer to fresh medium and therefore prolonged incubation without nutrient deficiency leads also to an increase of proliferating and fast-growing bacteria in liquid medium. Slow-growing organisms get the possibility to overcome their lag period and proliferate during longer incubation times. This re-diversification has a severe impact on results using liquid cultures, as several genera would be missed in case of earlier sampling and analysis. In this experiment, diversity stabilized between the third and fourth sampling time. The 16S rRNA gene data after 214 h of incubation revealed diverse individual cultures, stating that nitriles and their degradation products significantly impact bacterial community composition and diversity. In accordance with initial toxicity test, we could confirm an increased relative abundance of Gram-positive genera in cultures containing growth-suppressing nitriles such as phenylacetonitrile, 4-hydroxybenzonitrile and cyclohexanecarbonitrile. It appears that these organisms are more resistant to the negative influence of these specific compounds. Nevertheless, high abundance of *Paenibacillus* in the 4-hydroxybenzonitrile-containing sample was unexpected, as we could not find any confirmed nitrile-degrading enzyme for this genus in literature. However, putative nitrilase-encoding genes are present in publicly available *Paenibacillus* genomes. In addition, closely related cyanide dihydratases were just recently predicted for this genus (48). For *Brevundimonas*, also showing higher abundances in the nitrile-containing communities compared to the carboxylic acid-containing ones, database entries or literature on nitrile-degrading enzymes were not found. Nitrile degradation has been described for *Serratia* (49, 50), but potential nitrile-converting genes phylogenetically assigned to this genus were not predicted in the metagenome derived from the phenylacetonitrile-treated culture. Nevertheless, with an abundance of 40% in the respective culture, members of the *Serratia* genus seem to be well-adapted to phenylacetonitrile. Therefore, probably other mechanism lead to the nitrile tolerance. Thus, further investigation of all three genera could be of great value for the discovery of novel nitrile-degrading enzymes and resistance mechanisms.
A few reports mentioned nitrile-degrading activity of *Acinetobacter* (51, 52), but not for *Aeromonas*. The absence of putative nitrilases and nitrile hydratases for *Aeromonas* and only few nitrile hydratase subunits assigned to *Acinetobacter* indicate the rarity of respective genes in both genera. The reduced average abundance of both genera in nitrile-treated cultures is probably explained by a higher susceptibility to possible toxic effects of the nitriles and subsequent displacement of these genera by better adapted microorganisms such as *Paenibacillus* and *Brevundimonas*. *Pedobacter* is known for broad antibiotic resistance potential and heavy metal tolerance (53–55), but it seems like this genus is not adapted to nitrile-harboring or carboxylic acid-containing environments. In addition, genes encoding putative nitrilases or nitrile hydratases were not predicted to be derived from members of this genus. Interestingly, the same applies for putative nitrile-degrading enzymes derived from *Aneurinibacillus*, *Chryseobacterium*, and *Glutamicibacter*, but every genus shows growth in one nitrile-containing culture. Thus, the presence of so far unknown nitrile resistance or degradation mechanisms is indicated.

It is doubtful that with *Parastrongyloides* a parasite of Australian possums was living in German compost, but probably other related nematodes present in this environment could be able to survive in M9-HEPES, especially as *Parastrongyloides* shows growth in liquid medium (56). The assumption that nitriles and their handling are a common feature in nature (1–7) could be confirmed as many nitrile-degrading enzymes are commonly present in different genera. They could even be frequently identified in the control culture facing neither the nitrile itself nor the potential degradation product. Of special interest for further investigations could be the high abundance of nitrilase-encoding genes in the cyclohexanecarbonitrile-containing culture, demonstrating enrichment of respective genes. This could be an indicator for proliferation of a nitrile-degrading bacterium and consequently for active nitrile degradation.

Phylogenetic analysis revealed a high diversity of the encountered putative nitrilases as they were assigned to several different groups. The close clustering of aromatic nitrilases and ary lacetonitrilases was expected as both groups act on substrates containing aromatic structures. The *R. rhodochrous* K22 nitrilase preferably acts on aliphatic nitriles but is similar to benzonitrile-degrading nitrilases with respect to sequence (57). This could explain its clustering with other aromatic nitrilases, even though it has a different substrate specificity.
Conclusion and outlook. The here employed experimental setup for the investigation of the impact of nitriles and their degradation products on bacterial communities showed that these compounds significantly influence the composition of bacterial communities and partially lead to a strong reduction of diversity. In addition, it revealed strains with potential for remediation purposes such as Paenibacillus in case of 4-hydroxybenzonitrile.

In conclusion, this work provides a first insight on the influence of different nitriles and their corresponding carboxylic acids on microorganisms. A different susceptibility of Gram-negative (A. tumefaciens and E. coli) and Gram-positive (B. subtilis and C. glutamicum) test strains was observed during toxicity tests, but the number of tested organisms and chemicals should be increased to provide insights in the underlying mechanisms. The liquid culture experiment with a compost community as inoculum highlighted problems when transferring complex communities for several times to fresh medium, as the maximal optical densities were reduced each passage. Nevertheless, the analysis of the control cultures over time revealed a diversification and stabilization of the bacterial community composition, indicating that the time point for analysis of the chemically-treated cultures was appropriate. In addition, the toxic effects of certain nitriles (acetone cyanohydrin, 2-phenylpropionitrile, and pyruvonitrile) for microbial communities could be documented. Prevention of environmental contamination with these compounds seems therefore important, as bioremediation might be problematic. At the same time, lethal effects of the corresponding carboxylic acids were not observed, indicating specific toxicity of the nitriles.

The comparison of the cultures under the influence of nitriles with the cultures containing the respective carboxylic acids revealed many differences. The nitrile group exhibited a different effect on microorganisms than the acid counterpart. As some carboxylic acids show higher growth suppression and a strong influence on community composition, it seems reasonable to study the influence of educts and products of an enzymatic reaction. This should preferably be done before starting with enrichment or screening experiments for organisms harboring the corresponding enzymes to avoid unintended lethal effects. Especially putative nitrilases clustering with already characterized enzymes in the here presented phylogenetic tree are interesting candidates for further investigation, as chances for enzymatic activity are high. Additionally, a broader picture of nitrile influence on the nitrile-treated cultures could be gained by metatranscriptomic studies, as these have the potential to reveal stress responses, resistance mechanisms and novel degradation strategies. This knowledge could help developing novel or better biotransformation and bioremediation strategies and finally lead to a reduction of environmental contamination.
MATERIAL AND METHODS

**Media and stock solutions.** For growth of microorganisms, LB-Miller medium (10 g tryptone, 10 g NaCl and 5 g yeast extract per liter) (58) or M9-HEPES was used. M9-HEPES was prepared using 15.5 g HEPES, 1 g NH₄Cl and 0.5 g NaCl per liter. The pH was adjusted to 7.0 before autoclaving. Ten ml ATCC vitamin supplement (LGC Standards, Teddington, UK), 10 ml ATCC trace mineral supplement, 10 ml glucose solution (20 %), 1 ml MgSO₄ x 7H₂O solution (1 M), 1 ml CaCl₂ solution (14.7 g/l) and 1 ml thiamine-HCl solution (1 mg/ml) were added afterwards. For toxicity determination, following chemicals were solved in H₂O, DMSO, or ethanol and added to the medium: phenylacetonitrile (N1), succinonitrile (N2), acetonitrile (N3), crotononitrile (N4), 4-hydroxybenzonitrile (N5), acetone cyanohydrin (N6), cyclohexanecarbonitrile (N7), 2-phenylpropionitrile (N8), pyruvonic acid (N9), phenylacetic acid (A1), succinic acid (A2), acetic acid (A3), crotonic acid (A4), 4-hydroxybenzoic acid (A5), 2-hydroxyisobutyric acid (A6), cyclohexanecarboxylic acid (A7), 2-phenylpropionic acid (A8), and pyruvic acid (A9) (all chemicals from Sigma-Aldrich, Taufkirchen, Germany and TCI Chemicals, Eschborn, Germany). In case of carboxylic acids, an equimolar amount of ammonia was added in form of ammonia water (Carl Roth, Karlsruhe, Germany). For agar plates, 15 g agar per liter was added.

**Nitrile toxicity tests on selected strains.** Agrobacterium tumefaciens NTL4(pCF372)(pCF218) (59), Bacillus subtilis NCIB 3610 (60), Corynebacterium glutamicum ATCC 13032 (61) and Escherichia coli W3110 were used as model organisms during this study. Precultures of these strains were grown in LB medium at 30 °C and 180 rpm (Multitron shaker, Infors, Einsbach, Germany) for 24 h. Afterwards, each strain was plated on a quadrant of a LB or M9-HEPES plate containing one of the nine nitriles or the respective carboxylic acid in a concentration of 5, 10, 15, 20 or 25 mM. As control, plates with LB or M9-HEPES were used. Every 24 h, the growth on the plates was analyzed and categorized on a scale of 0 to 4 by comparison with the control. In case of no growth a 0 was given, whereas similar growth compared to the control was treated as 4. The remaining values were given for approx. 25% (=1), 50% (=2) and 75% (=3) of colony size and opaqueness when compared to the control.

**Nitrile toxicity on an environmental community.** Compost (100 g, pH 7.5) of the Experimental Botanical Garden Göttingen, Germany (51°33'22.6"N 9°57'16.2"E) were solved in 500 ml ddH₂O and subsequently filtered with a 2.7 µm GF/D glass fiber filters (Whatman, Little Chalfont, UK). Baffled flasks (total volume 100 ml) were filled with 35 ml M9-HEPES containing one of the nitriles or carboxylic acids in the above-mentioned tested concentrations (Table 1). Flasks filled only with M9-HEPES served as control. All conditions were prepared in triplicates. The solved and filtered compost (5 ml) were added to each baffled flask and the cultures were incubated at 25 °C and 75 rpm (Multitron shaker). To monitor the growth of the cultures, the optical density at 600 nm (OD\textsubscript{600}) was measured. Two prevent starvation of the microorganisms, cultures were transferred after three days to fresh medium.
containing the same nitrile or carboxylic acid at the concentrations used before. The new cultures were adjusted to an OD<sub>600</sub> of 0.1 and incubated again at 25 °C and 75 rpm. In case that cultures showed no growth, the cultures were not transferred to fresh medium but further incubated. The procedure was repeated two additional times after two days of incubation per growth period. The final cultures were again incubated for two days.

**DNA extraction.** DNA of the compost sample solved in water was isolated before and after filtering using the PowerMax soil DNA isolation kit (MO BIO Laboratories, Carlsbad, CA, USA) according to the manufacturer’s protocol. DNA of the M9-HEPES-containing enrichment cultures was isolated using the MasterPure Complete DNA and RNA purification kit as recommended by the manufacturer (Epicentre, Madison, WI, USA). Concentration of isolated DNA was measured using a NanoDrop ND-1000 (Thermo Fisher Scientific, Waltham, MA, USA) or with a Quant-iT dsDNA HS assay kit and a Qubit fluorometer as recommended by the manufacturer (Invitrogen, Carlsbad, CA, USA).

**16S rRNA gene sequencing.** For amplification of the V3-V4 region of bacterial 16S rRNA genes, the primers S-D-Bact-0341-b-S-17 and S-D-Bact-0785-a-A-21 (62) were used. The primers contained adapters for Illumina MiSeq sequencing (Illumina, San Diego, CA, USA). The DNA isolated from solved compost or M9-HEPES-containing enrichment cultures was used as template. PCR reaction mixtures (50 µl final volume) contained 10 µl 5-fold Phusion GC buffer, 0.2 µM of each of the primers, 0.2 µM MgCl<sub>2</sub>, 2.5 µl DMSO, 200 µM of each of the four dNTPs, 1 U of Phusion DNA polymerase (Thermo Fisher Scientific), and 25 ng template DNA. For amplification, the following protocol was used: initial denaturation at 98°C for 1 min, 25 cycles of denaturation at 98 °C for 45 s, annealing at 60 °C for 45 s, and elongation at 72 °C for 30 s, followed by final elongation at 72 °C for 5 min. PCR reactions for each culture were performed in triplicate, pooled in equal amounts, and purified with the NucleoMag 96 PCR kit (Macherey-Nagel, Düren, Germany) according to the manufacturer’s protocol. For quantification of the PCR products, a Qubit fluorometer was used in combination with the Quant-iT dsDNA HS assay kit as recommended by the manufacturer (Invitrogen). The purified amplicons were indexed using the Nextera XT DNA library prep kit according to the instructions of the manufacturer (Illumina). Dual index paired-end sequencing (2 x 300 bp) with v3 chemistry was done on an Illumina MiSeq platform as recommended by the manufacturer (Illumina).

**16S rRNA gene sequence analysis.** CASAVA data analysis software (Illumina) was used for demultiplexing and clipping of sequence adapters from raw sequences. Before removing sequences with an average quality score below 20 and unresolved bases with split_libraries_fastq.py from QIIME 1.9.1 (63), paired-end sequences were merged using PEAR v0.9.10 with default parameters (64). Default settings of cutadapt 1.14 (65) were used for removal of non-clipped reverse and forward primer sequences. Subsequently, the UNOISE2 pipeline of USEARCH 9.2.64 (66) was used to remove,
dereplicate, and denoise reads shorter than 380 bp, leading to zero-radius operational taxonomic units (zOTUs). Furthermore, chimeric sequences were removed. For this purpose, UCHIME2 was used in reference mode against the SILVA SSU database version 128 (67). To create a zOTU table, the quality-filtered sequences were mapped to chimera-free OTUs using USEARCH. With parallel_assign_taxonomy_blast.py taxonomic classification of the zOTU sequences against the SILVA database was done. Filter_otu_table.py was used for removal of chloroplasts, unclassified OTUs, and extrinsic domain OTUs. Finally, the lowest number of sequences by random subsampling (12,100 reads per sample) was used for sample comparison at the same surveying effort.

**Metagenome sequencing and analysis.** DNA isolated from each culture triplicate was pooled in equal amounts. Paired-end metagenome sequencing was performed with an Illumina HiSeq 4000 (2 × 150 bp) using the Nextera XT DNA library prep kit. For quality-trimming and verification of the paired-end reads, Trimmomatic version 0.36 (68) and FastQC version 0.11.5 (69) were used. Assembly and coverage calculation was performed with SPAdes version 3.10 (70). Contigs >200 bp were annotated with Prokka 1.11 (71). For taxonomic assignment of nitrilases, a BLASTn search (72) of the nitrile containing contigs was performed against the NCBI non-redundant database. Abundance of nitrile-degrading genes was calculated by mapping quality-filtered reads against the sequence-based identified putative nitrile-converting genes with Bowtie2 version 2.3.4.2 (73). Normalization was done by dividing the number of mapped nucleotides by the total number of nucleotides. Subsequently, these values were normalized against the number of putative target genes.

**Phylogenetic tree.** Evolutionary analyses were conducted in MEGA X version 10.0.5 (74). Alignment of the amino acid sequences was done with MUSCLE (75) and standard settings. The evolutionary history was inferred by using the Maximum Likelihood method and LG model (76) as recommended by build-in model test tool of MEGA X. The tree with the highest log likelihood (-31012.19) is shown. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using a JTT model, and then selecting the topology with superior log likelihood value. A discrete Gamma distribution was used to model evolutionary rate differences among sites (5 categories (+G, parameter = 1.1643)). This analysis involved 276 amino acid sequences. A total of 491 positions were in the final dataset. Extensive subtree pruning regrafting and a very weak branch swap filter were used. Phylogeny was tested by calculating 500 bootstrap replicates and nodes with values below 50% were condensed for the final image. Reference sequences were recovered from the NCBI database (accession numbers in parentheses): For aromatic nitrilases, *Rhodococcus rhodochrous* J1 (BAA01994) (77), *R. ruber* NHB-2 (CCN27135) (78), and *Gordonia terrae* MTCC 8139 (AGR86048) (79) were used as references. Aliphatic nitrilases were derived from *Comamonas testosteroni* (AAA82085) (80), *Synechocystis* sp. PCC6803 (BAA10717) (81),
and *R. rhodochrous* K22 (BAA02127) (82), while arylacetonitrilases were obtained from *Alcaligenes faecalis* JM3 (BAA02684) (83), *Pseudomonas fluorescens* EBC191 (AAW79573) (84), and *Pseudomonas putida* MTCC 5110 (ABV21758) (85). β-Cyano-L-alanine nitrilases were derived from *P. fluorescens* Pf0-1 (ABA74312) (86, 87), *Pseudomonas protegens* Pf-5 (AAY92181) (86, 88), and *Pseudomonas pseudoalcaligenes* CECT 5344 (CDM40486) (89, 90). Cyanide dihydratases were retrieved from *Pseudomonas stutzeri* AK61 (AIY29195) (91), *Bacillus pumilus* C1 (AF392815) (92), and *B. pumilus* 8A3 (AAN77003) (92). Furthermore, three amidases were included, originating from *Pseudomonas aeruginosa* PAC142 (P11436) (93), *R. rhodochrous* M8 (AAX83004) (94), and *Geobacillus pallidus* RAPc8 (AAO23013) (95).

**Accession numbers.** Raw data of 16S rRNA gene analysis and metagenome sequences as well as assembled metagenomes have been deposited at NCBI under the following biosample accession numbers: compost sample, SAMN09932763; filtered compost sample, SAMN09932764; control after 68 h, SAMN09932785; control after 188 h, SAMN09932808; control after 166 h, SAMN09932809; control after 214 h, SAMN09932848; phenylacetonitrile enrichment, SAMN09930795; succinonitrile enrichment, SAMN09932325; acetonitrile enrichment, SAMN09932597; crotononitrile enrichment; SAMN09932603; 4-hydroxybenzonitrile enrichment, SAMN09932608; cyclohexanecarbonitrile enrichment, SAMN09932617; phenylacetic acid enrichment, SAMN09932620; succinic acid enrichment, SAMN09932700; acetic acid enrichment, SAMN09932745; crotonic acid enrichment, SAMN09932756; 4-hydroxybenzoic acid enrichment, SAMN09932757; cyclohexanecarboxylic acid enrichment, SAMN09932758.

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3.2. Nitrile-degrading bacteria isolated from compost

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Supplementary data can be found on the enclosed DVD

Author contributions

Performed the experiments: RE
Analyzed the data: RE, DS
Wrote the publication: RE, DS, RH, RD
Conceived the study: RH, RD
Nitrile-Degrading Bacteria Isolated from Compost

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Keywords: acetonitrile, nitrilase, nitrile hydratase, amidase, Flavobacterium, Pseudomonas, Rhodococcus, Variovorax

INTRODUCTION

Nitriles are a diverse group of organic compounds with –C≡N as functional group. Most nitriles are slightly cytotoxic but some cause severe toxic effects. More than 120 naturally occurring nitriles without considering cyanogenic glycosides are present in terrestrial and marine habitats, especially in plant components such as almonds or other fruit pits. The most common group of naturally occurring nitriles are cyanogenic glycosides, which can be found in more than 100 plant families as well as in fungi, bacteria, and animals. This group of molecules can be chemically or enzymatically hydrolyzed, leading to the release of highly toxic hydrogen cyanide and thereby act as natural defense compound (Fleming, 1999). For detoxification, two enzymatic pathways for the degradation of nitriles are known. The first one involves nitrilases (EC 3.5.5.1), a subgroup of the carbon-nitrogen hydrolase superfamily, which degrade nitriles directly to carboxylic acids and ammonia. The second one is a bi-enzymatic pathway using nitrile hydratases (NHases; EC 4.2.1.84) for the degradation of nitriles to amides and amidases (EC 3.5.1.4) for the subsequent degradation to carboxylic acids and ammonia (Gong et al., 2012). The enzymatic hydrolysis of nitriles proceeds under mild reaction conditions, whereas the chemical hydrolysis is dependent on acidic or alkaline conditions and high temperatures. The latter also results in the production of large quantities of byproducts and inorganic waste (Clouthier and Pelletier, 2012; Vergne-Vaxelaire et al., 2013).

Consequently, nitrile-converting enzymes are of increasing industrial importance with respect to green chemistry. A constantly increasing number of nitrile-derived amides [e.g., acrylamides or carboxylic acids (e.g., glycolic acid)] are produced with these enzymes (Schmid et al., 2001; Panova et al., 2007). In addition, nitrilases can be used for the treatment of nitrile-polluted wastewater (Li et al., 2016) and other environmentally-friendly bioremediation processes (Gong et al., 2012).

Here, we report data on the taxonomic composition of an enrichment culture with acetonitrile as nitrogen source. In addition, we present eight individual bacterial draft genome sequences of isolates obtained from this enrichment. The genome content of these isolates was analyzed with respect to genes responsible for the nitrile-degrading phenotype. Genome and average nucleotide identity analysis indicated that the isolated bacterial strains are affiliated to the species Rhodococcus erythropolis, Flavobacterium sp., Variovorax boronicumulans, Pseudomonas sp., and Pseudomonas kilonensis.

MATERIALS AND METHODS

Isolation of the Bacteria

Compost (100 g, pH 7.5) of the Experimental Botanical Garden Göttingen, Germany (51°33′22.6″N 9°57′16.2″E) was suspended in 500 ml H₂O and filtered with a 2.7 μm GF/D glass fiber filter (Whatman, Little Chalfont, UK). Enrichment and control cultures were each initiated with 750 μl of the resulting filtrate. Enrichment cultures were grown in 40 ml M9 medium
(Atlas, 2010) with ATCC trace mineral supplement (LGC Standards, Teddington, UK), 10 mM glucose as carbon source and 25 mM acetonitrile as sole nitrogen source at 25°C for 4 days. Control cultures were grown in same medium but the medium contained 18.7 mM NH₄Cl instead of acetonitrile. The enrichment culture was streaked on solid M9 medium with 25 mM acetonitrile as sole nitrogen source and incubated at 25°C. The obtained 33 colonies showed four distinct colony morphologies, and were picked and purified over five rounds of incubation. Growth in the absence of an added nitrogen source was not recorded.

16S rRNA Gene Sequencing

Genomic DNA of the liquid cultures and the specific isolates was extracted using the MasterPure complete DNA and RNA purification kit (Epicentre, Madison, WI, USA), while DNA isolation of the compost sample was done with the PowerMax soil DNA isolation kit (MO BIO Laboratories, Carlsbad, CA, USA). The bacterial composition was determined by amplicon-based analysis of the V3-V4 region of the 16S rRNA gene using the bacterial primers S-D-Bact-0341-b-S-17 and S-D-Bact-0785-a-A-21 (Klindworth et al., 2013) with adapters for Illumina MiSeq sequencing (Illumina, San Diego, CA, USA). The PCR reaction mixture (50 µl) contained 10 µ1 5-fold Phusion GC buffer, 200 µM of each of the four dNTPs, 2.5 µ1 DMSO, 0.2 µM of each primer, 200 µM MgCl₂, 1 U of Phusion polymerase (Thermo Fisher Scientific, Waltham, MA, USA), and 25 ng of isolated DNA as template. The following cycling scheme was used: initial denaturation at 98°C for 1 min and 25 cycles of denaturation at 98°C for 45 s, annealing at 60°C for 45 s, and extension at 72°C for 30 s, followed by a final extension at 72°C for 5 min. PCR reactions were performed in triplicate for each sample. The resulting PCR products were pooled in equal amounts and purified using the NucleoMag 96 PCR kit (Macherey-Nagel, Düren, Germany) as recommended by the manufacturer. Quantification of the PCR products was performed using the Quant-iT dsDNA HS assay kit and a Qubit fluorometer as recommended by the manufacturer (Invitrogen, Carlsbad, CA, USA). Indexing of the PCR products was performed with Nextera XT DNA library prep kit as described by the supplier (Illumina). Sequencing was performed with the Illumina MiSeq platform using the dual index paired-end approach (2 × 300 bp) and v3 chemistry.

The 16S rRNA genes of specific isolates were amplified with the primer pair 08f (5’-AGAGTTTGATCCTGGCTC-3’) and 1504r (5’-TACCTTGTTACGACTT-3’). The previously mentioned cycling scheme was modified to an annealing temperature of 40°C and an extension time of 45 s. Sanger sequencing of the PCR products was done by Seqlab (Göttingen, Germany).

16S rRNA Gene Amplicon Analysis

Demultiplexing and clipping of sequence adapters from raw amplicon sequences were performed by employing CASAVA data analysis software (Illumina). Paired-end sequences were merged using PEAR v0.9.10 with default parameters (Zhang et al., 2014). Subsequently, sequences with an average quality score lower than 20 and containing unresolved bases were removed with the split_libraries_fastq.py script of QIIME 1.9.1 (Caporaso et al., 2010). Non-clipped reverse and forward primer sequences were removed by employing cutadapt 1.12 with default settings (Martin, 2011). USEARCH version 9.2.64 was used following the UNOISE pipeline (Edgar, 2010). In detail, reads shorter than 380 bp were removed, dereplicated, and denoised with the UNOISE2 algorithm of USEARCH resulting in zero-radius operational taxonomic units (zOTUs). Additionally, chimeric sequences were removed using UCHIME2 in reference mode against the SILVA SSU database release 128 (Yilmaz et al., 2014). All quality-filtered sequences were mapped to chimera-free OTUs and a zOTU table was created using USEARCH. Taxonomic classification of the picked reference sequences (zOTUs) was performed with parallel_assign_taxonomy_blast.py against the same SILVA database. Extrinsic domain OTUs, chloroplasts, and unclassified OTUs were removed from the dataset by employing filter_otu_table.py. Sample comparisons were performed at same surveying effort, utilizing the lowest number of sequences by random subsampling (31,000 reads per sample).

RESULTS AND DISCUSSION

Enrichment of the compost sample with M9 minimal medium containing acetonitrile as sole nitrogen source resulted in microbial growth, indicating the presence of acetonitrile-degrading organisms. The bacterial diversity and composition of the compost sample and the enrichment culture were determined via amplicon-based 16S rRNA gene analysis. The compost sample used as starting material for enrichment comprised 520 zOTUs, which were reduced to 63 during enrichment (Supplementary Table S1). The dominant bacterial genera after
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<th>Isolate</th>
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<th>GC content (%)</th>
<th>No of Genes</th>
<th>No of tRNAs</th>
<th>No of rRNAs</th>
<th>Protein, no. of amino acids (focus tag)</th>
<th>Closest similar protein, no. of amino acids (accession no. of similar protein)</th>
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<th>Region of similar amino acids (%)</th>
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enrichment were *Aeromonas* (33%) and *Pseudomonas* (55%) with similar abundances in the control sample (Figure 1A). In the enrichment culture the relative abundances of *Flavobacterium* and *Bacillus* were 5 and 2%, respectively, and in the control culture 2 and 0.4%, respectively. *Chryseobacterium*, *Paenibacillus*, *Pedobacter*, and *Sphingobacterium* varied between 1.1 and 1.3% in the control culture, but showed relative abundances below 0.1% in the acetonitrile-containing enrichment. In contrast, *Variovorax* (3%) was only detected in the acetonitrile enrichment culture in significant amounts. We performed isolation experiments from enrichment cultures on solid media with acetonitrile as sole nitrogen source. A total of 33 isolates (Figure 1B) were recovered of which eight isolates (ACN1 to ACN8, Table 1) showed a different 16S rRNA gene sequence. All isolates were able to grow with acetonitrile as sole nitrogen source in axenic culture. No growth was observed in the absence of an added nitrogen source. The genomes of all eight isolates were sequenced and analyzed with the focus on genes potentially involved in nitrile-degradation (Table 1). Based on average nucleotide identity, tetra-nucleotide signatures and in silico DNA-DNA hybridization, the eight isolates were affiliated to the genera *Flavobacterium*, *Pseudomonas*, *Rhodococcus* and *Variovorax*. Besides *Rhodococcus* all isolated strains belonged to genera detected during 16S rRNA gene analysis of the bacterial community in the enrichment culture.

**Rhodococcus (ACN1)**

In contrast to all other isolates, only slight traces of *Rhodococcus* could be found during 16S rRNA gene analysis. Nevertheless, this isolate was obtained multiple times from the enrichment, indicating its ability to degrade acetonitrile. The reason for the discrepancy between 16S rRNA gene analysis of the community and the isolation results may be due to non-optimal media conditions for *Rhodococcus* in the enrichment culture, leading to low abundance. Genome sequencing of *R. erythropolis* ACN1 and quality-filtering resulted in 2,641,652 paired-end reads resulting in a draft genome of 7.24 Mbp with a 93.9-fold coverage. Similarity searches for putative genes involved in nitrile-degradation revealed two genes (BKP42_65670 and BKP42_65680), coding for one of the two putative NHase subunits. These enzymes are known to be responsible for nitrile degradation in various *R. erythropolis* strains (Kaufmann et al., 1999; Brandão et al., 2003; Vejvoda et al., 2007; Kamble et al., 2013). The deduced protein sequences showed highest identity to the alpha subunit of *Rhodococcus* sp. N-774 (Ichhata et al., 1989) and the beta subunit of *R. erythropolis* deep-sea strain 122-AN065 (Brandão et al., 2003). The deduced acetamidases (BKP42_40660 and BKP42_26750) are most similar to acetamidases of *R. erythropolis* and *Rhodococcus* sp. 311R (Ehsani et al., 2015). The two aliphatic amidases of the isolated strain (BKP42_54900 and BKP42_64640) most resemble an acylamid amidohydrolase from *Gordonia desulfuricans* and an amidase of *Rhodococcus* sp. R312 (Fournand et al., 1998). An additional amidase (BKP42_65660) is most similar to an amidase of *Rhodococcus* sp. N-774 (Table 1).

**Variovorax (ACN3)**

Members of the genus *Variovorax* were prime candidates for active nitrile degradation, as the genus appeared only in significant abundances in the bacterial community after enrichment with acetonitrile. With the isolation of ACN3 identified as *V. boronicumulans* a member of this genus was recovered. Sequencing and assembly of 3,471,160 paired-end reads resulted in a draft genome of 7.14 Mbp (131-fold coverage). Although *V. boronicumulans* is a species described just recently (Miwa et al., 2008), evidence for nitrile degradation by this microorganism was reported before (Zhang et al., 2012; Liu et al., 2013). A search for genes encoding putative nitrile-degrading enzymes resulted in the identification of genes for two possible degradation pathways, the first one via a nitrilase (BKP43_17560) and the second one via an NHase (BKP43_58000 and BKP43_58010) and amidase (BKP43_28100). The most similar enzymes to the nitrilase, alpha and beta subunits of NHase, and the aliphatic amidase are an amidohydrolase, NHase subunits and acylamide amidohydrolase of another *V. boronicumulans* strain, respectively (Table 1).

**Pseudomonas (ACN4, ACN5, ACN7, and ACN8)**

With a relative abundance of over 50%, *Pseudomonas* is the major genus of the bacterial community in the acetonitrile enrichment culture. Several *Pseudomonas* species such as *P. chlororaphis* and *P. fluorescens* are able to use nitriles as nitrogen source.
and harbor genes encoding nitrilases, or NHases and amidases (Nagasawa et al., 1987; Kiziak et al., 2005; Howden et al., 2009). The presence of these genes could be due to the plant habitat of some Pseudomonas species, which is rich in different and unusual nitriles like indole-3-carboxyl nitrile (Rajniak et al., 2015).

Isolates ACN4 and ACN7 were affiliated to P. kilonensis, a species described in 2001 (Sikorski et al., 2001). Sequencing of the ACN4 genome resulted in 2,239,346 paired-end reads, an average coverage of 96.7-fold, and a draft genome of 6.55 Mbp. Genes encoding a putative nitrilase (BSF43_29400) and amidas (BSF43_35090) were detected, which showed highest identity to a Pseudomonas nitrilase and a putative acylamide amidohydrolase of P. sp. Root401 (Bai et al., 2015), respectively.

The genome assembly of isolate ACN7 was based on 2,049,404 paired-end reads and resulted in a draft genome of 6.49 Mbp and a 74.9-fold coverage. The GC-content is 61.0%. The draft genome encodes a nitrilase (BSG18_13010) and an amidase (BSG18_29450) similar to a nitrilase of P. sp. Ep R1 (Chiellini et al., 2014) and an acylamide amidohydrolase of P. sp. Root401, respectively.

The two Pseudomonas isolates ACN5 and ACN8 could not be assigned to a specific Pseudomonas species. The genome of ACN5 (6.69 Mbp, 61.3-fold coverage) was assembled from 1,350,588 paired-end reads. Potential genes for two nitrilases, one NHase and two amidases were predicted in the ACN5 genome. The aliphatic nitrilase (BSF40_48900) showed highest similarity to a nitrilase of Pseudomonas sp. UW4 (Duan et al., 2013), while the other nitrilase (BSF40_42250) most resembles a nitrilase of Pseudomonas sp. GM48 (Brown et al., 2012). The NHase alpha and beta subunits (BSF40_24660 and BSF40_24650) are related to the corresponding ones of Pseudomonas and P. sp. UW4. The predicted amidases (BSF40_24670 and BSF40_59650) are similar to an amidase of P. sp. GM33 and GM49 (Brown et al., 2012). For the assembly of the ACN8 genome, 1,505,826 paired-end reads were used, resulting in a draft genome of 6.34 Mbp and 66.0-fold coverage. Genes for two nitrilases (BSF44_34690 and BSF44_52950) were predicted, which resemble two different nitrilases from P. fluorescens. Furthermore, two genes encoding putative amidases (BSF44_36240 and BSF44_50390) similar to an amidase of P. composti and P. sp. GM74 (Brown et al., 2012) were detected.

Flavobacterium (ACN2 and ACN6)

The two isolates ACN2 and ACN6 from the acetonitrile enrichment culture were affiliated to the Gram-negative genus Flavobacterium. A total of 1,503,824 paired-end reads were used for the assembly of the ACN2 genome, resulting in a draft genome of 5.40 Mbp with an average coverage of 72.7-fold. ACN6 genome assembly of 2,409,950 paired-end reads yielded a draft genome of 5.13 Mbp with average coverage of 120.4-fold. Genes encoding putative nitrile-degrading enzymes were not detected in both draft genomes. As the nitrile degrading capacity of both strains was independently verified multiple times by growth experiments with acetonitrile as sole nitrogen source, the presence of so far unknown genes and pathways for nitrile utilization is indicated. In addition, the ability to overcome the nitrogen limitation via nitrogen fixation from air was experimentally excluded as no growth could be monitored in liquid M9 medium without any added nitrogen source. Furthermore, typical genes for nitrogen fixation such as dinitrogenases and dinitrogenase reductases were not identified in the genome sequences. Thus, two explanations are possible for the discrepancy between the observed phenotype of both strains and the lack of putative genes for nitrile degradation. First, genes responsible for nitrile degradation could not be annotated as they are located in contig gaps, as both genomes are still in the draft state. This is unlikely as most contig gaps are due to repetitive regions longer than the read length of the used sequencing technology (Whiteford et al., 2005; Chaisson et al., 2015). Second, genes for new types of nitrile degradation enzymes are present. The only other available study on nitrile degradation by Flavobacteria reports weak degradation of 3-cyanopyridine by F. aquatile IFO 3772, F. suaveolens IFO 3752 and F. rigense IAM 1238 (Kato et al., 2000), but genes responsible for the observed phenotype were not reported and genomes of the three strains are not available. Thus, further analyses are required to unravel the basis of the nitrile-degrading phenotype of the flavobacterial isolates.

CONCLUSION

The here presented data revealed the potential of specifically designed enrichment experiments for the isolation of organisms with a desired metabolic activity like nitrile degradation. Our study provided new candidates to cover the industrial demand for new nitrile-degrading biocatalysts as part of a green chemistry. While for the isolates R. erythropolis ACN1, V. boronicumulans ACN2, P. kilonensis ACN4 and ACN7 as well as P. sp. ACN5 and ACN8 nitrilases and/or NHases and amidases could be annotated, the nitrile-degrading pathway of F. sp ACN2 and ACN6 remains to be unraveled.

ACCESSION NUMBERS

The draft genome sequences of all eight organisms have been deposited at GenBank with the following accession numbers: R. erythropolis ACN1, MRCL00000000; F. sp. ACN2, MRCM00000000; V. boronicumulans ACN3, MRCN00000000; P. kilonensis ACN4, MRCO00000000; P. sp. ACN5, MRCP00000000; F. sp. ACN6, MRCQ00000000; P. kilonensis ACN7, MRCR00000000; P. sp. ACN8, MRC50000000. Raw sequence data of all genomes are available at the NCBI SRA archive and linked to the respective BioSamples. The sequences of the 16S rRNA gene analysis are linked to the BioSamples SAMN07278921 (inoculum), SAMN0278947 (control) and SAMN07278950 (acetonitrile enrichment). Strains have been submitted to the DSMZ German Collection of Microorganism and Cell Cultures and are available on request.

AUTHOR CONTRIBUTIONS

RD and RH conceived the study. RE performed the experiments. RE and DS analyzed the data. All authors interpreted the results.
wrote the manuscript and reviewed the final version of the manuscript.

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REFERENCES


Egelkamp et al. Nitrile-Degrading Bacteria Isolated from Compost

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fenvs.2017.00056/ full#supplementary-material


Egelkamp, et al. Nitrile-Degrading Bacteria Isolated from Compost


**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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3.3. From sequence to function:
A new workflow for nitrilase identification

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*RE and IF contributed equally to this work

Supplementary data can be found on the enclosed DVD

Author contributions

Performed the experiments: IF
Analyzed the data: \textbf{RE}, IF
Wrote the publication: \textbf{RE}, IF, RH, RD
Conceived the study: \textbf{RE}, RH, RD
ABSTRACT

This study describes a workflow for simple and fast acquisition of nitrilase candidates from large metagenomic sequence sets. For identification of active enzymes, a NADH-coupled high-throughput assay was established. Purification of enzymes could be omitted as crude extract containing the overexpressed putative nitrilases was used. In addition, long incubation times could be avoided by combination of nitrile and NADH conversion in a single reaction. This allowed the direct measurement of nitrile degradation and provides not only insights into substrate specificity but also in degradation efficiency. The novel assay was subsequently used for investigation of seventy putative nitrilases previously identified during sequence-based screens of metagenomes derived from nitrile-treated microbial communities. Six enzymes showed activity and the most interesting nitrilase was purified and analyzed. The activity of the characterized arylacetonitrilase exhibited a broad pH range and a high long-term stability. The enzyme showed high specificity for phenylacetonitrile with $K_m$ of 1.29 mM and $V_{max}$ of 13.85 U/mg protein. In conclusion, we provide a setup for simple and rapid analysis of putative nitrilases from sequence to function, demonstrated by the identification, isolation, and characterization of the here described arylacetonitrilase.

INTRODUCTION

Nitriles are often toxic compounds with $-\text{C≡N}$ as functional group. They are widespread in nature and present in plants in form of cyanoglycosides (1). Other naturally occurring nitriles are e.g. cyanohydrins in fungi and arthropods, different antibiotics in bacteria, or ricine and phenylacetonitrile in plants (2). For the enzymatic degradation of nitriles, two pathways are known. The first is the direct conversion of nitriles to corresponding carboxylic acids via nitrilases (EC 3.5.5.1) and the second the degradation of nitriles to corresponding amides via nitrile hydratases (NHases) (EC 4.2.1.84) and the subsequent hydrolysis of amides to carboxylic acids and ammonia using amidases (EC 3.5.1.4). Nitrile degrading catalysts can be found in many organisms including bacteria, filamentous fungi, yeasts, and plants (3–5). In industry, these enzymes are used for production of bulk chemicals such as nicotinic acid or glycolic acid (6, 7). Identification of novel nitrile-degrading enzymes is important due to their industrial use and the often observed substrate and product inhibition under production conditions (8–11). Many nitrile-degrading enzymes were found via (meta)genome mining techniques. A functional screening approach could be applied to identify nitrilase genes from plasmid libraries containing metagenomic DNA from different sources (12–14). In silico screening of (meta)genomic sequence data is used to identify putative nitrilases and nitrile hydratases in publicly available sequence space (15). Although sequence-based approaches lead very quickly to many new enzyme candidates, they remain to be predictions until functional verification. To overcome this limitation and take advantage of the
constantly growing metagenomic sequence pool, procedures are needed for efficient verification of nitrilase activity.

The focus of this investigation was to establish a workflow which enables fast functional verification of nitrilases identified by sequence-based screenings. As starting material, six metagenomes which were obtained during determination of nitrile-influence on microbial communities (Chapter 3.1) were chosen. A procedure was established to reduce the 70 putative nitrilase-encoding genes derived from sequence-driven mining of the metagenomes to candidates with verified enzyme activity. Serving as prove of concept, one enzyme candidate was heterologously expressed and purified. Subsequently, the recovered enzyme and its activity were characterized including the determination of optimal pH, temperature range, relative stability, and substrate specificity. In addition, influence of divalent ions and other substances on enzyme activity, and enzyme kinetics were determined.

RESULTS

Screening optimization. Most modern high-throughput nitrilase assays have drawbacks such as low sensitivity or work only under specific conditions. One of the most promising approaches is a NADH-coupled assay in which ammonia released by the nitrilase reaction is consumed by a glutamate dehydrogenase (GDH) under conversion of NADH to NAD⁺. The conversion of this cofactor is monitored at 340 nm and reveals activity of the tested (putative) nitrilases (15). However, it requires extended incubation times for nitrilase reaction and provides only qualitative information and no insights regarding enzymatic efficiency. We modified the assay to allow fast and simple screening without additional incubation steps in a single reaction mixture. Therefore, concentration of all test ingredients was optimized to allow continuous measurement of NADH conversion in a 96-well plate, leading to first insights into efficiency of nitrile degradation. In comparison to the previous method, the amount of NADH was increased to 0.5 mM for a larger measuring range and the concentration of α-ketoglutarate was reduced to prevent inhibition of the GDH reaction. These modifications were tested by the use of an already characterized nitrilase from *Rhodococcus rhodochrous* K22 (16, 17) (Figure 1).

To reduce identification of false positives or enzymes with low activity, a threshold was defined. During the first 15 minutes of measurement, total conversion of succinonitrile, fumaronitrile, and crotononitrile by the K22 nitrilase could be observed. As activity for these substrates was described before (16, 17), only reactions taking place in the first 15 minutes were considered as nitrilase activity during subsequent experiments.
Figure 1: Nitrile degradation of the *R. rhodochrous* K22 nitrilase. The assay was performed in triplicate at 37 °C in a 96-well plate. NADH conversion was monitored at 340 nm. Fumaronitrile, succinonitrile, and crotononitrile were degraded after 15 min, thereby defining the threshold (red line) for the experimental time.

**Analysis of putative nitrilases.** Six metagenomes of microbial communities obtained from phenylacetonitrile-, succinonitrile-, acetonitrile-, crotononitrile-, 4-hydroxybenzonitrile-, or cyclohexanecarbonitrile-containing cultures (Chapter 3.1) served as source for putative nitrilases. In total, 70 annotated putative nitrilases were identified in these metagenomes and subsequently analyzed (Data Set S1). Initially, all enzyme candidates were compared with 46 reference nitrilases obtained from SWISS-Prot to relate the deduced protein sequences to already characterized enzymes. Results led to 13 protein clusters (Data Set S2) and reduced our data set to 60 unique candidates. The first three clusters contained characterized as well as putative nitrilases ranging from 5.96 to 38.68 kDa, 12.7 to 40.2 kDa, and 10.41 to 38.62 kDa. Six clusters consisted just of known and characterized nitrilases, whereas the remaining four clusters included just putative enzyme sequences. As the smallest characterized prokaryotic nitrilase (from *Pyrococcus abyssi* GE5) consists of 262 amino acids (18), all putative nitrilases smaller than this were checked for ribosomal binding sites (RBS) to exclude artificial peptides caused by incomplete metagenomic assembly. Finally, 37 candidates were selected for cloning and overexpression in *E. coli*.

After amplification of the respective genes from metagenomic DNA and sequence verification, frameshift mutations were observed in some cases and led to the exclusion of these candidates. In other cases, point mutations were observed, which were included into subsequent analyses as not every point mutation leads to non-functional proteins. Finally, a total number of 26 putative nitrilases were sub-cloned (Data Set S1). Subsequent overexpression and verification via SDS page revealed 13 putative nitrilases.
The previously established assay was used to verify nitrilase activity with the 13 putative nitrilase candidates using nine different nitriles as substrate (Table 1). Six of the putative enzymes showed nitrilase activity for at least one nitrile. Most were active with succinonitrile or fumaronitrile. In addition, one converted the aromatic phenylacetonitrile (Figure 2). Thus, the workflow presented here has shown a way to successfully narrow down bioinformatically identified enzyme candidates to the interesting functional representatives.

**Table 1: Overexpressed putative nitrilases.** Metagenome: metagenomic origin of the genes. Variant: Variants obtained by point mutations (for sequences, see Data Set S1). Origin: gene sequences were used for a blastn search against the NCBI nr database and best hits are shown; Q: query cover; E: e value; I: ident; A: accession. Substrate: substrate determined with high-throughput assay. SUN: succinonitrile; CAN: acetonitrile; CRN: crotononitrile; CCN: cyclohexanecarbonitrile; PAN: phenylacetonitrile; FUN: fumaronitrile.

<table>
<thead>
<tr>
<th>Metagenome</th>
<th>Nitrilase</th>
<th>Variant</th>
<th>Mass [kDa]</th>
<th>Origin</th>
<th>Substrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>SUN</td>
<td>D3C71_601570</td>
<td>B</td>
<td>36.12</td>
<td><em>Variovorax boronicumulans</em> (Q: 100%, E: 0.0, I: 98%, A: KY937903)</td>
<td>PAN</td>
</tr>
<tr>
<td>SUN</td>
<td>D3C71_775930</td>
<td>B</td>
<td>33.0</td>
<td><em>Pseudomonas fluorescens</em> (Q: 99%, E: 0.0, I: 87%, A: CP010945)</td>
<td>-</td>
</tr>
<tr>
<td>SUN</td>
<td>D3C71_946180</td>
<td>C</td>
<td>35.13</td>
<td><em>Pseudomonas sp.</em> (Q: 99%, E: 0.0, I: 94%, A: CP003880)</td>
<td>SUN, FUN</td>
</tr>
<tr>
<td>ACN</td>
<td>D3C72_113100</td>
<td>A</td>
<td>38.7</td>
<td><em>Janthinobacterium sp.</em> (Q: 99%, E: 0.0, I: 87%, A: CP000269)</td>
<td>-</td>
</tr>
<tr>
<td>ACN</td>
<td>D3C72_306600</td>
<td>A</td>
<td>38.62</td>
<td><em>Pseudomonas sp.</em> (Q: 97%, E: 0.0, I: 89%, A: AP017423)</td>
<td>SUN, FUN</td>
</tr>
<tr>
<td>ACN</td>
<td>D3C72_447890</td>
<td>A</td>
<td>32.93</td>
<td><em>Pseudomonas fluorescens</em> (Q: 99%, E: 0.0, I: 87%, A: CP010945)</td>
<td>-</td>
</tr>
<tr>
<td>ACN</td>
<td>D3C72_810700</td>
<td>C</td>
<td>36.74</td>
<td><em>Cupriavidus basilensis</em> (Q: 98%, E: 0.0, I: 87%, A: CP010537)</td>
<td>SUN, FUN</td>
</tr>
<tr>
<td>CRN</td>
<td>D3C73_384910</td>
<td>A</td>
<td>35.2</td>
<td><em>Pseudomonas sp.</em> (Q: 100%, E: 0.0, I: 98%, A: CP003880)</td>
<td>SUN, FUN</td>
</tr>
<tr>
<td>CRN</td>
<td>D3C73_384910</td>
<td>B</td>
<td>35.18</td>
<td><em>Pseudomonas sp.</em> (Q: 100%, E: 0.0, I: 98%, A: CP003880)</td>
<td>SUN, FUN</td>
</tr>
<tr>
<td>CRN</td>
<td>D3C73_406050</td>
<td>A</td>
<td>32.84</td>
<td><em>Pseudomonas sp.</em> (Q: 100%, E: 0.0, I: 95%, A: CP003880)</td>
<td>-</td>
</tr>
<tr>
<td>CRN</td>
<td>D3C73_406050</td>
<td>B</td>
<td>32.86</td>
<td><em>Pseudomonas sp.</em> (Q: 100%, E: 0.0, I: 91%, A: CP003880)</td>
<td>FUN</td>
</tr>
<tr>
<td>CRN</td>
<td>D3C73_579180</td>
<td>G</td>
<td>32.91</td>
<td><em>Pseudomonas sp.</em> (Q: 100%, E: 0.0, I: 97%, A: CP003880)</td>
<td>-</td>
</tr>
<tr>
<td>CCN</td>
<td>D3C75_459950</td>
<td>B</td>
<td>32.83</td>
<td><em>Pseudomonas sp.</em> (Q: 100%, E: 0.0, I: 91%, A: CP003880)</td>
<td>-</td>
</tr>
</tbody>
</table>
Figure 2: Substrate specificity of enzyme D3C71_601570. The assay was performed at 37 °C in a 96-well plate. NADH conversion was monitored at 340 nm. Degradation of phenylacetonitrile was observed until the 15 min threshold (red line).

Characterization of enzyme D3C71_601570. To prove the previous observations, we chose the phenylacetonitrile-degrading enzyme candidate for detailed characterization and verification of enzymatic activity under defined conditions. For further experimental characterization, a sequence encoding a His$_6$ tag was added to the N-terminal end of the gene. Subsequently, the enzyme was produced and purified. SDS-PAGE analysis of the enzyme preparation confirmed the from the sequence deduced molecular mass of 36 kDa. The purified enzyme was subjected to a detailed characterization with focus on optimal enzymatic conditions, stability, and kinetics. Phenylacetonitrile was used as standard substrate, as the nitrilase showed only activity for this compound during high-throughput screening.

Investigations regarding pH dependency showed pH 6 to be optimal, whereby at pH 5.5 to 8 about 90% enzyme activity could still be observed. The limits of the enzyme were observed at pH 5 and 10.5, as only 20% activity remained (Figure 3A). Optimum temperature of the enzyme was determined in the range of 25 °C to 60 °C at pH 6. Highest activity was measured between 45 °C and 55 °C, with an optimum at 50 °C (Figure 3B). At 60 °C, nitrilase activity was reduced to 30%. Furthermore, stability of the nitrilase was tested over several weeks while the enzyme was stored at 4 °C. Remaining catalytic activity >80% could be observed during the first three weeks, followed by a slow decline (Figure 3C). After three months, 8% of the initial activity remained.
Results

Figure 3: Temperature and pH dependence, and long-term stability of enzyme D3C71_601570. (A) pH optimum. Reactions were run for 2 min at 37 °C in the respective buffers (0.1 M) containing 1.25 µg purified protein and 5 mM phenylacetonitrile. 100% relative activity corresponds to 2.79 U/mg. (B) Temperature optimum. Reactions were run for 2 min at various temperatures using citrate-phosphate buffer (pH 6.0), 1.25 µg purified protein, and 5 mM phenylacetonitrile. 100% relative activity corresponds to 5.68 U/mg. (C) Stability assay. Enzyme was stored at 4 °C in a buffered system (50 mM monosodium phosphate, 300 mM sodium chloride, 250 mM imidazole, pH 8.0). Reactions were run for 2 min at 50 °C using citrate-phosphate buffer (pH 6.0), 1.25 µg purified protein, and 5 mM phenylacetonitrile. 100% relative activity corresponds to 5.68 U/mg.

Catalytic activity of the nitrilase was tested with the substrates also used for the high-throughput assay, verifying its specificity for phenylacetonitrile under the tested conditions. To determine activity of the enzyme for structurally related compounds, four aromatic nitriles (2-phenylbutyronitrile, 3-phenylpropionitrile, benzonitrile, and mandelonitrile) were additionally tested. With 2-phenylbutyronitrile and benzonitrile as substrate, nitrilase activity was not detected but the enzyme exhibited reduced activity with mandelonitrile and 3-phenylpropionitrile (22 and 16 % of activity measured with phenylacetonitrile, respectively) (Table 2). In conclusion, the metagenome-derived nitrilase D3C71_601570 is highly specific for phenylacetonitrile and belongs to the group of arylacetonitrilases.

To assess the effect of potential inhibitors and solvents on enzyme activity, the purified enzyme was tested against different chemicals. For detection of essential thiol residues, the thiol-binding reagents HgCl₂ and AgNO₃ were applied to the reaction mixture. The observed decline of enzyme activity to 1.6 and 2.0%, respectively, indicated that the functionality of nitrilase D3C71_601570 depends on these residue (Table 3). As the metal-chelating agent EDTA had no effect on enzymatic function, dependence on metal-cofactors could be excluded. This was further supported by missing effects of divalent ions such as Mg²⁺, Fe²⁺, and Mn²⁺ on nitrilase activity. Interestingly, Zn²⁺ increased activity to 114%. Most likely, stimulating effect of Zn²⁺ derives from a different mechanism, i.e. increased protein stability in the presence of Zn²⁺. Furthermore, treatment with H₂O₂ and SDS led to expected reduction of activity, which is most likely due to denaturation of the nitrilase.
Strong inhibitory effects were encountered in the presence of DMSO, acetone, and chloroform even at 5% concentration (Table 4). At this concentration, nitrilase activity was less sensitive to methanol, ethanol, isopropanol, and glycerol treatment. At 20% concentration, only glycerol did not abolish the main enzymatic activity.

Table 2: Substrate specificity of enzyme D3C71_601570. Reactions were run for 15 min at 50 °C using citrate-phosphate buffer (pH 6.0), 1.25 µg purified protein, and 20 mM nitrile. Activity with phenylacetonitrile as substrate (5.19 U/mg) was set as 100% relative activity.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Relative activity [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenylacetonitrile</td>
<td>100.00 ± 1.19</td>
</tr>
<tr>
<td>Succinonitrile</td>
<td>0.17 ± 0.03</td>
</tr>
<tr>
<td>Acetonitrile</td>
<td>0.00 ± 0.01</td>
</tr>
<tr>
<td>Crotononitrile</td>
<td>0.08 ± 0.01</td>
</tr>
<tr>
<td>4-Hydroxybenzonitrile</td>
<td>0 ± 0.00</td>
</tr>
<tr>
<td>Acetone cyanohydrin</td>
<td>0.13 ± 0.02</td>
</tr>
<tr>
<td>Cyclohexanecarbonitrile</td>
<td>0.09 ± 0.01</td>
</tr>
<tr>
<td>Fumaronitrile</td>
<td>0.45 ± 0.04</td>
</tr>
<tr>
<td>2-Phenylpropionitrile</td>
<td>0.52 ± 0.05</td>
</tr>
<tr>
<td>Mandelonitrile</td>
<td>22.01 ± 0.61</td>
</tr>
<tr>
<td>2-Phenylbutyronitrile</td>
<td>0.07 ± 0.01</td>
</tr>
<tr>
<td>3-Phenylpropionitrile</td>
<td>15.81 ± 0.80</td>
</tr>
<tr>
<td>Benzonitrile</td>
<td>0.48 ± 0.32</td>
</tr>
</tbody>
</table>

Table 3: Effects of various compounds on the activity of enzyme D3C71_601570. Reactions were run for 2 min at 50 °C using citrate-phosphate buffer (pH 6.0), 1.25 µg purified protein, 5 mM phenylacetonitrile, and 1 mM putative inhibitor. Enzyme activity with phenylacetonitrile as substrate without additions was set as 100% relative activity (5.68 U/mg).

<table>
<thead>
<tr>
<th>Compound</th>
<th>Relative activity [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>EDTA</td>
<td>104.45 ± 1.62</td>
</tr>
<tr>
<td>DTT</td>
<td>76.21 ± 3.29</td>
</tr>
<tr>
<td>H2O2</td>
<td>24.56 ± 2.35</td>
</tr>
<tr>
<td>HgCl2</td>
<td>1.57 ± 0.36</td>
</tr>
<tr>
<td>AgNO3</td>
<td>1.96 ± 0.18</td>
</tr>
<tr>
<td>CaCl2</td>
<td>106.29 ± 2.71</td>
</tr>
<tr>
<td>MnSO4</td>
<td>98.01 ± 3.80</td>
</tr>
<tr>
<td>MgSO4</td>
<td>101.87 ± 5.92</td>
</tr>
<tr>
<td>FeSO4</td>
<td>98.25 ± 5.65</td>
</tr>
<tr>
<td>CuSO4</td>
<td>86.98 ± 3.82</td>
</tr>
<tr>
<td>CoSO4</td>
<td>81.01 ± 2.60</td>
</tr>
<tr>
<td>ZnCl2</td>
<td>114.00 ± 2.04</td>
</tr>
<tr>
<td>Sodium azide</td>
<td>98.89 ± 2.13</td>
</tr>
<tr>
<td>SDS</td>
<td>12.02 ± 2.77</td>
</tr>
</tbody>
</table>
Table 4: Effect of different organic solvents on activity of enzyme D3C71_601570. Reactions were run for 2 min at 50 °C using citrate-phosphate buffer (pH 6.0), 1.25 µg purified protein, and 5 mM phenylacetonitrile. Enzyme activity with phenylacetonitrile as substrate without additions was set as 100% relative activity (5.68 U/mg).

<table>
<thead>
<tr>
<th>Solvent</th>
<th>Relative activity of enzyme with 5% v/v organic solvent [%]</th>
<th>Relative activity of enzyme with 20% v/v organic solvent [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methanol</td>
<td>88.01 ± 6.18</td>
<td>9.26 ± 1.97</td>
</tr>
<tr>
<td>Ethanol</td>
<td>85.22 ± 7.12</td>
<td>4.82 ± 0.41</td>
</tr>
<tr>
<td>Glycerol</td>
<td>81.11 ± 8.68</td>
<td>76.65 ± 9.06</td>
</tr>
<tr>
<td>Isopropanol</td>
<td>77.55 ± 9.43</td>
<td>3.63 ± 0.91</td>
</tr>
<tr>
<td>DMSO</td>
<td>4.80 ± 1.06</td>
<td>2.32 ± 0.72</td>
</tr>
<tr>
<td>Acetone</td>
<td>28.96 ± 4.79</td>
<td>0.14 ± 0.07</td>
</tr>
<tr>
<td>Chloroform</td>
<td>24.81 ± 3.44</td>
<td>3.70 ± 1.38</td>
</tr>
<tr>
<td>Tolul</td>
<td>8.12 ± 2.13</td>
<td>2.44 ± 1.60</td>
</tr>
</tbody>
</table>

Finally, kinetic constants of nitrilase D3C71_601570 were determined. The reaction rate increased with substrate concentration until a plateau was reached at 5 mM. Between 10 and 50 mM a rapid decrease in activity was recorded, followed by a slow decline until 100 mM (Figure 4A), indicating inhibition by the substrate. Based on these data, kinetic constants were calculated with a non-linear model due to superior precision compared to linear methods like Lineweaver-Burk plots (19). Phenylacetonitrile was converted at 37 °C with a $K_M$ of 1.29 mM and $V_{max}$ of 13.85 U/mg, reaching the saturation point at 6 mM (Figure 4B). The enzyme is highly specific for this nitrile and the low $K_M$ value indicated high affinity for the substrate.

Figure 4: Influence of substrate concentrations on the reaction rate of enzyme D3C71_601570. Reactions were run with phenylacetonitrile as substrate at 37°C in a UV/Vis spectrophotometer. A: steady-state measurement until 100 mM substrate concentration. B: steady-state measurement until saturation point at 6 mM substrate concentration. C: Lineweaver-Burk plot.
DISCUSSION

High-throughput nitrilase screening. In this study, a fast and simple high-throughput assay for screening of functional nitrilases was established. In comparison to the previously described method (15), it improves the total screening time by combining nitrilase reaction and subsequent ammonia conversion in a single reaction. Furthermore, it allows constant measurement of nitrile degradation and provides insights into the efficiency of the tested enzymes. Unfortunately, the measurement is limited by the GDH as only nitrilase reactions slower than the GDH can be monitored in detail. In addition, different enzymes in the crude extract can lead to ammonia release and influence the measurement. Other enzymes may also be involved in the reverse reaction of NAD\(^+\) to NADH, leading to distorted results. To minimize the effect of these factors, additional precautions have to be taken. Most important is the use of crude extract containing the empty expression vector as negative control. To further reduce the likelihood of false positive hits, a time limit for nitrilase reaction can be used. Based on the reactions observed with the well-characterized _R. rhodochrous_ K22 nitrilase (16, 17), this threshold was set at 15 minutes.

After establishment of screening parameters, 70 putative nitrilases identified in metagenomes of nitrile-treated microbial communities were taken as input. Combining the sequence-based nitrilase clusters obtained during bioinformatical analysis with results from the high-throughput assay, only candidates clustering with characterized nitrilases showed enzymatic activity. Skipping clusters consisting just of putative nitrilases during future screenings would reduce subsequent work but involves the risk of missing nitrilases with novel sequences. Therefore, decisions on the cost-benefit ratio of including these clusters in further analyses must be individually made.

Although only preselected putative nitrilase genes were amplified, not all of them could be overexpressed or the gene products formed inclusion bodies in _E. coli_. These results could be due to different codon usage of _E. coli_ TOP10 compared to the native hosts of the enzymes. Diverging rare codons in _E. coli_ could lead to suboptimal expression and low amounts of the desired protein or unwanted mRNA degradation (20–23). Furthermore, misfolding and aggregation due to missing translational pauses could occur (24). One possibility to avoid these problems is using lower temperatures during protein expression or the utilization of low-copy-number plasmids (25–28). In addition, codon-optimization of the candidate genes for _E. coli_ could be performed. However, these methods were not applied as they are mostly not suitable for large numbers of candidates.

Characterization of an arylacetonitrilase. The prediction and subsequent isolation of a phenylacetonitrile-degrading nitrilase shows the reliability of the novel nitrilase high-throughput screening. According to sequence similarity, the isolated arylacetonitrilase is affiliated to _Variovorax_
boronicumulans (Table 1). Nitrile-degrading abilities of this species have been described before, but phenylacetonitrile degradation has not been mentioned (29–31).

With 36 kDa the here described enzyme is close to molecular mass of other known phenylacetonitrile-degrading nitrilases such as the nitrilases from *Synechocystis* sp. PCC 6803 (40 kDa), *Bradyrhizobium japonicum* USDA 110 (34.5 kDa), or *Alcaligenes faecalis* ATCC 8750 (36 kDa) (32–34). In contrast, the broad pH range of D3C71_601570 activity is not that common (34, 35). In addition, it exhibits an unusual long-term stability, as similar nitrilases lose up to 80% activity in ten days (32, 35), whereas the here characterized enzyme still retained 80% activity after three weeks.

One major difference of the here presented enzyme to all other phenylacetonitrile-degrading nitrilases is its narrow substrate range. Phenylacetonitrile was the preferred substrate and even small differences in molecular structure seem to inhibit proper catalytic function. During high-throughput assay, minor degradation of other nitriles was observed in the crude extract of the strain producing enzyme D3C71_601570. None of these substrates were degraded by the purified enzyme, underlining again the importance of the 15-minute threshold to avoid identification of false positive candidates. Nevertheless, the differences can also come from the His<sub>6</sub>-tag added for purification (36, 37).

CONCLUSION

In this study, we could show that a combination of targeted sequence data analyses with a high-throughput activity assay can assist in the fast identification of candidates for functional nitrilases. Due to real-time measurement of NADH degradation, the here presented assay does not only provide hints on possible substrates but also on the catalytic efficiency of tested enzymes. Furthermore, the use of crude extract is less time consuming compared to methods demanding purified enzymes.

One of the active enzymes showed phenylacetonitrile-degrading abilities. The most likely to *Variovorax* belonging nitrilase is highly specific for this arylacetonitrile, as closely related molecules were barely degraded. To our knowledge, the enzyme shows unique properties, as other nitrilases usually degrade several (related) substrates. Therefore, it can give new insights into sequence function relationship of arylacetonitrilases and help to develop better biocatalysts.

MATERIAL AND METHODS

Metagenomic DNA and putative nitrilases. Metagenomic DNA originated from soil enrichment cultures containing either phenylacetonitrile, succinonitrile, acetonitrile, crotononitrile, 4-hydroxybenzonitrile, or cyclohexanecarbonitrile (Chapter 3.1). Genbank accession numbers of the annotated metagenomes are as follows: phenylacetonitrile, RCUE00000000; succinonitrile, RCUP00000000; acetonitrile, RCUQ00000000, crotononitrile, RCUN00000000; 4-hydroxybenzonitrile, RCUF00000000; cyclohexanecarbonitrile, RCUO00000000.
**Bioinformatic analysis.** Nitrilase reference data were obtained from the SWISS-Prot database (38) (date of search: December 12th, 2017). The data were further processed by removing misannotated enzymes, mere subunits, or nitrile hydratases designated as nitrilases. Clustering of data was done with the web suite CD-HIT (39). A fasta file containing the SWISS-Prot nitrilases and the putative nitrilases was used as input and a sequence identity cut-off of 40% was set for clustering.

**Nitrilase positive control.** An already characterized aliphatic nitrilase from *Rhodococcus rhodochrous* K22 was used as positive control for establishing the high-throughput nitrilase assay (16, 17). The nitrilase was codon-optimized for *Escherichia coli* K12 derivates using the web suite JCat (http://www.jcat.de) (40) and synthesized by Integrated DNA Technologies (Leuven, Belgium).

**Media.** LB-Miller (10 g tryptone, 10 g NaCl, and 5 g yeast extract per liter) (41) was used for growth of microorganisms. For agar plates, 15 g agar per liter was added. Media were sterilized by autoclaving at 121 °C for 20 min.

**PCR protocol.** 50 µl PCR reaction mixture contained 10 µl 5-fold Phusion HF buffer, 200 µM of each dNTP, 0.2 µM of both primers (Data Set S3), 3% DMSO, 50 ng metagenomic DNA as template, 1 U of Phusion polymerase (Thermo Fisher Scientific, Waltham, MA, USA) and was filled up with ddH2O. Initial denaturation was done at 98 °C for 5 min, followed by 30 cycles of denaturation (98 °C, 30 s), annealing (temperature based on primer properties, 30 s) and elongation (72 °C, 45 s/kbp). The final elongation was for 5 min at 72 °C.

**Plasmids.** Amplified putative nitrilase-encoding genes (Data Set S2) were sub-cloned into the pBAD18 vector system (42). For selection, a final concentration of 100 µg/ml ampicillin was used. The fidelity of the constructs was checked by sequencing (Microsynth Seqlab, Göttingen, Germany) with forward primer 5’-ATTAGCGGATCCTACCTGACG-3’ and reverse primer 5’-CAGACCGCTTCTGCGTTCTG-3’.

**Strains and transformation.** Chemically competent *Escherichia coli* TOP10 cells (Thermo Fisher Scientific, Waltham, MA, USA) were transformed according to the manufacturer’s protocol. Subsequently, cells were plated on ampicillin-containing LB plates and incubated overnight at 37 °C.

**Heterologous expression.** *E. coli* TOP10 containing the pBAD18 vector with desired insert was grown in LB medium at 37 °C overnight at 180 rpm (Innova 44 shaker, New Brunswick Scientific, Nürtingen, Germany). The preculture was used to inoculate 10 ml LB medium to an OD600 of 0.1, which was subsequently incubated for 1.5 h at 37 °C and 180 rpm (Innova 44 shaker) to reach an OD600 of 0.6–0.8. For induction of expression, 1.5% L-(+)-arabinose were added, followed by 6 h of incubation.
Cell disruption and purification of His$_6$-tagged protein. Cells containing the overexpressed (His$_6$-tagged) nitrilase were washed twice with 1x LEW buffer from the Protino Ni-TED kit (Macherey-Nagel, Düren, Germany) and resuspended in 1.5 ml buffer with 40 µg/ml DNase I and 0.1% w/v lysozyme. The cells were then disrupted at least three times by French press at $1.38 \times 10^8$ Pa (Thermo Fisher Scientific). The soluble fractions of the thereby recovered solution were obtained by centrifugation at 6000 $\times$ g and 4 °C for 20 min to remove cell debris. For purification, the resulting supernatant was then loaded according to the manufacturer’s protocol. The purified enzyme was further analyzed by 12% SDS-PAGE and protein concentration was determined using Bradford method (43) with bovine serum albumin as standard.

Nitriles. Nitriles were solved at a concentration of 2.7 M in DMF (Merck KGaA, Darmstadt, Germany). These stock solutions were filtered sterile and stored at 4 °C. Following nitriles were used: phenylacetonitrile, acetonitrile (all TCI Deutschland GmbH, Eschborn, Germany), succinonitrile, crotonitrile, 4-hydroxybenzonitrile, acetone cyanohydrin, cyclohexanecarbonitrile, fumaronitrile, and 2-phenylpropionitrile (all Sigma-Aldrich Chemie GmbH, Munich, Germany). In addition, mandelonitrile, 2-phenylbutyronitrile, benzonitrile (all Sigma-Aldrich), and 3-phenylpropionitrile (Alfa Aesar, Haverhill, MA, USA) were used for characterization of the arylacetonitrilase.

High-throughput nitrilase assay. Degradation of nitriles was measured by monitoring the release of ammonia in a coupled enzymatic reaction (44). The ammonia reacts with NADH and α-ketoglutarate through a glutamate dehydrogenase (GDH) to glutamate. The reduced cofactor is monitored by its absorbance change at 340 nm. For screening of putative nitrilases, the high-throughput assay of Vergne-Vaxelaire et al. (15) was adjusted to allow real-time measurements without time-consuming incubation steps (15). The assay was performed with technical replicates in a flat-based 96-well plate. The reaction mixture (final volume 250 µl) contained 0.5 mM NADH, 1 mM α-ketoglutarate, 37 µl of a 2.7 M nitrile stock solution (solved in DMF), 20 µg crude extract (added at the end), and 1 U/ml GDH from bovine liver type II (Sigma-Aldrich Chemie GmbH) and 50 mM Tris-HCl (pH 8.0) buffer. As negative control, crude extract containing the pBAD18 cloning vector was used. The reaction mixture was incubated at 37 °C in a Synergy 2 microplate reader (BioTek Instruments GmbH, Bad Friedrichshall, Germany). The absorbance was constantly measured for 1.5 h at 340 nm. For calculation, the values of the pBAD18 crude extract (negative control) were subtracted from that of the nitrilase crude extract. To avoid identification of false positives, only reactions completed during the first 15 minutes of measurement were considered as nitrilase activity.
Nitrilase characterization. The selected nitrilase (NCBI accession D3C71_601570) was characterized using a colorimetric assay for detection of ammonia (45). Nitrilase reaction was performed in a 500 µl mixture containing 25 µl nitrile (100 mM, solved in DMF), 1.25 µg purified His6-tagged nitrilase and 0.1 M citrate-phosphate buffer (pH 6.0) for 2 min at 50 °C. The experimental environment was at 23 °C. 111 µl of this reaction were transferred into 222 µl sodium phenate, followed by 333 µl of 0.01% sodium nitroprusside and 333 µl of hypochlorite. The reaction mixture was incubated for 15 min at 27 °C in darkness and subsequently absorption was measured at 630 nm. Same procedure was used for substrate determination with 20 mM of the respective nitrile and 15 min incubation.

Nitrilase inhibitors. Effect of various compounds on nitrilase activity was tested. Incubation was carried out at 50 °C for 2 min in the standard reaction mixture containing a putative inhibitor at 1 mM or 5% v/v and 20% v/v when testing organic solvents.

Nitrilase stability. For stability testing, the His6-tagged purified enzyme was stored in Protino® Ni-TED elution buffer (Macherey-Nagel, Düren, Germany) containing 1 mM sodium azid at 4 °C in the dark.

Steady-state activity assay. The steady-state kinetics for arylacetonitrilase activities were measured using a GDH (bovine liver type II) to detect the concomitant oxidation of NADH at 340 nm and 37 °C using a UV/Vis spectrophotometer (Cary 100 UV-Vis, Varian Medical Systems, Darmstadt, Germany). The reaction mixture contained 0.5 mM NADH, 1 mM α-ketoglutarate, 0.1 mM to 100 mM of phenylacetonitrile (final volume 37 µl, nitrile solved in DMF), 5 µg purified nitrilase (added at the end), and 1 U/ml GDH from bovine liver type II (Sigma-Aldrich Chemie GmbH) filled up to 1000 µl with 0.1 M HEPES (pH 8.0) buffer. Negative control did not contain the nitrilase. The slope (ΔA₃₄₀) was determined by Cary WinUV version 3.0 (Varian Medical Systems). The $k_{cat}$ values were plotted against phenylacetonitrile concentrations. Further, the macroscopic constants $k_{cat}$ and $K_M$ were analyzed using the Michaelis-Menten equation of R package ‘drc’ (46).

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REFERENCES


3.4. Novel approach for identification of biocatalysts

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Supplementary data can be found on the enclosed DVD

Author contributions

Performed the experiments: RE
Analyzed the data: RE, HK
Wrote the publication: RE, RH, RD
Conceived the study: RE, RH, RD
INTRODUCTION

Molecules containing a \(-\text{C≡N}\) functional group are known as nitriles. These structurally diverse and often toxic compounds are widespread in nature and present mainly in plants (1). Enzymatic degradation of nitriles is performed by two different known pathways: the direct degradation of nitriles to corresponding carboxylic acids and ammonia by nitrilases, or the hydration of nitriles to amides by nitrile hydratases (NHases) and the subsequent degradation to carboxylic acids and ammonia by amidases. In addition to their natural occurrence, nitriles are also an important compound in industry for the production of different bulk chemicals and pharmaceuticals (2–5). Due to the increasing interest in green chemistry and therefore a high demand on nitrile-degrading enzymes, identification of novel nitrilases is of increasing importance (6).

In general, screening for novel enzymes by (meta)genomic approaches is done employing two methods. The first is a function-based screening using (meta)genomic libraries and their expression in heterologous hosts like *Escherichia coli* (7). This method could lead to the discovery of genes encoding unknown enzymes but requires high effort. Furthermore, heterologous protein expression could lead to non-functional proteins and therefore prevent their detection. The second approach is a sequence-based approach in which (meta)genomic datasets are screened for genes of interest (8). This method is limited to homologues of already known enzymes, making the detection of truly novel proteins impossible. In addition, genes merely identified by sequence similarities do not necessarily exhibit the targeted function.

During the last years, a third screening approach based on differential transcriptomics arose (9–11). For this purpose, an organism is treated under at least two different conditions and respective transcriptomes are sequenced. By statistical analysis, up- and downregulated genes of each treatment are identified, providing insight into genes important for specific conditions. Drawback of this method are the complex models for normalization and the high computational power needed for analysis (12, 13).

In this study, an experimental setup for the (meta)transcriptome-based discovery of genes encoding nitrile-degrading enzymes is presented. For microbial growth, media with acetonitrile and for comparison with acetic acid and ammonia as sole carbon and nitrogen sources were used. Therefore, genes upregulated in acetonitrile-treated cultures compared to acetic acid-containing samples are promising candidates for novel nitrile-degrading enzymes. In addition, unknown transporters or resistance mechanisms can be identified by this method. In a previous experiment, *Rhodococcus erythropolis* ACN1 was isolated from compost and its acetonitrile-degrading ability experimentally confirmed (14). The isolate was used during the here presented study to establish growth conditions
and bioinformatic analysis strategies. Subsequently, the approach was applied to a complex microbial community for identification of genes encoding novel nitrilases by differential metatranscriptomics.

**RESULTS**

**Establishment of a transcriptome-based nitrilase screening.** *R. erythropolis* ACN1 was grown in minimal medium containing acetic acid and ammonia (ACA) or acetonitrile (ACN) as sole carbon and nitrogen sources. Growth of the ACN cultures was delayed by approx. 25 hours when compared with ACA samples. After initial lag phase, similar growth pattern was observed (Figure 1). Final optical density (OD₆₀₀) of the ACA and the ACN cultures were almost identical (1.6 and 1.5, respectively).

![Figure 1: Growth of *R. erythropolis* ACN1 under different treatments.](image)

For both conditions, transcriptomes were harvested during logarithmic growth. After quality-filtering, lowest number of transcriptome reads was 17.8 million and more than 99% of the filtered reads mapped onto the genomic sequence of *R. erythropolis* ACN1 (14). Subsequent normalization and analysis were performed with DESeq2, an algorithm optimized for transcriptome comparison (15). The algorithm offers a false discovery rate (FDR) cutoff to reduce number of erroneously identified differentially expressed genes. To minimize number of false positives, a cutoff of 0.025 was chosen for further analysis. With this setting, 186 differentially expressed genes were identified of which 26 were upregulated and 160 downregulated in the ACN culture when compared to the ACA treatment.

Besides this statistical parameter, the difference between both conditions considered as significantly up- or downregulated was defined. For that purpose, a logarithmic fold change (log₂FC) between the normalized gene read counts was calculated. As a log₂FC of 2.0 is commonly used and streamlines subsequent analysis (16–18), respective values were applied in this study. This second criterium reduced the number of differentially expressed candidate genes to 23 of which 4 were upregulated...
and 19 downregulated in the ACN culture (Table 1). Further analysis focused on upregulated genes, as they are most promising to comprise the targeted genes encoding nitrile-degrading enzymes. The upregulated genes belonged to two groups. The first group contained an amidase (BKP42\_54900) with a log\(_2\)FC of 2.1, while the second group consisted of the remaining upregulated genes and was in proximity to the amidase. All members were upregulated at least six-fold, indicating a strong effect of the nitrile on their expression. Nevertheless, none of the respective annotations indicated a nitrile-related function, as they consisted of a tubulin-like protein (BKP42\_54850), a hypothetical protein (BKP42\_54860), and a GTPase domain (BKP42\_54870). When performing additional blastn and blastp searches against the NCBI nr database, annotation of the aliphatic amidase was confirmed, whereas best hits for the remaining upregulated genes always consisted of hypothetical proteins (Table 2). An additional motif search against the NCBI CDD (conserved domain database) supported annotation of the amidase and the tubulin family GTPase.

Table 1: Significantly up- or downregulated genes in *R. erythropolis* ACN1 when grown with acetonitrile compared to treatment with acetic acid and ammonia. Differential expression was analyzed with DESeq2 with an FDR of 0.025 and log\(_2\)FC of ≥2 as cutoff. Positive values (bold) represent genes upregulated under ACN treatment; negative values represent downregulated genes.

<table>
<thead>
<tr>
<th>Locustag</th>
<th>Gene</th>
<th>log(_2)FC</th>
</tr>
</thead>
<tbody>
<tr>
<td>BKP42_13460</td>
<td>CAAX amino terminal protease self-immunity</td>
<td>-2.0</td>
</tr>
<tr>
<td>BKP42_14580</td>
<td>putative siderophore transport system permease protein YfiZ precursor</td>
<td>-2.2</td>
</tr>
<tr>
<td>BKP42_22340</td>
<td>hypothetical protein</td>
<td>-2.2</td>
</tr>
<tr>
<td>BKP42_29290</td>
<td>Enterobactin exporter EntS</td>
<td>-2.3</td>
</tr>
<tr>
<td>BKP42_29300</td>
<td>Ferrienterobactin-binding periplasmic protein precursor</td>
<td>-2.3</td>
</tr>
<tr>
<td>BKP42_29310</td>
<td>Linear gramicidin synthase subunit B</td>
<td>-2.6</td>
</tr>
<tr>
<td>BKP42_29320</td>
<td>Flavodoxin</td>
<td>-2.3</td>
</tr>
<tr>
<td>BKP42_29330</td>
<td>Isochorismatase</td>
<td>-2.6</td>
</tr>
<tr>
<td>BKP42_29340</td>
<td>2,3-dihydroxybenzoate-AMP ligase</td>
<td>-2.4</td>
</tr>
<tr>
<td>BKP42_29350</td>
<td>2,3-dihydro-2,3-dihydroxybenzoate dehydrogenase</td>
<td>-2.0</td>
</tr>
<tr>
<td>BKP42_29370</td>
<td>Isochorismate synthase DhbC</td>
<td>-2.0</td>
</tr>
<tr>
<td>BKP42_29390</td>
<td>hypothetical protein</td>
<td>-3.3</td>
</tr>
<tr>
<td>BKP42_39400</td>
<td>Ferrienterobactin-binding periplasmic protein precursor</td>
<td>-3.4</td>
</tr>
<tr>
<td>BKP42_39410</td>
<td>Enterochelin esterase</td>
<td>-2.3</td>
</tr>
<tr>
<td>BKP42_43800</td>
<td>putative siderophore transport system permease</td>
<td>-2.0</td>
</tr>
<tr>
<td>BKP42_43810</td>
<td>Ferric enterobactin transport system permease protein FepG</td>
<td>-2.7</td>
</tr>
<tr>
<td>BKP42_54850</td>
<td>Tubulin-like protein CetZ</td>
<td>3.2</td>
</tr>
<tr>
<td>BKP42_54860</td>
<td>hypothetical protein</td>
<td>3.9</td>
</tr>
<tr>
<td>BKP42_54870</td>
<td>Tubulin/FtsZ family, GTPase domain</td>
<td>3.9</td>
</tr>
<tr>
<td>BKP42_54900</td>
<td>Aliphatic amidase</td>
<td>2.1</td>
</tr>
<tr>
<td>BKP42_65440</td>
<td>Fe(3+)–citrate-binding protein YfmC precursor</td>
<td>-2.1</td>
</tr>
<tr>
<td>BKP42_65450</td>
<td>Iron(3+)–hydroxamate-binding protein FhuD precursor</td>
<td>-2.0</td>
</tr>
<tr>
<td>BKP42_65460</td>
<td>putative siderophore transport system ATP-binding protein YusV</td>
<td>-2.1</td>
</tr>
</tbody>
</table>
Table 2: Best blastn, blastp, and NCBI CDD hits for significantly upregulated genes of *R. erythropolis* ACN1 when grown with acetonitrile compared to treatment with acetic acid and ammonia. Q: query cover; E: e value; I: ident; A: accession.

<table>
<thead>
<tr>
<th>Locustag</th>
<th>Annotation</th>
<th>Blastn</th>
<th>Blastp</th>
<th>CDD</th>
</tr>
</thead>
<tbody>
<tr>
<td>BKP42_54850</td>
<td>Tubulin-like protein CetZ</td>
<td><em>Rhodococcus</em> sp. NJ530 plasmid unnamed 4 (Q: 100%, E: 0.0, I: 99%, A: CP034156)</td>
<td>Hypothetical protein [<em>Rhodococcus erythropolis</em>](Q: 100%, E: 0.0, I: 100%, A: WP_095972695)</td>
<td>-</td>
</tr>
<tr>
<td>BKP42_54860</td>
<td>Hypothetical protein</td>
<td><em>Rhodococcus</em> sp. NJ530 plasmid unnamed 4 (Q: 100%, E: 7e-177, I: 99%, A: CP034156)</td>
<td>Hypothetical protein [<em>Rhodococcus erythropolis</em>](Q: 100%, E: 7e-80, I: 100%, A: PBI91134)</td>
<td>-</td>
</tr>
<tr>
<td>BKP42_54870</td>
<td>Tubulin/FtsZ family, GTPase domain</td>
<td><em>Rhodococcus</em> sp. NJ530 plasmid unnamed 4 (Q: 100%, E: 0.0, I: 99%, A: CP034156)</td>
<td>Hypothetical protein [<em>Rhodococcus erythropolis</em>](Q: 100%, E: 0.0, I: 100%, A: WP_095972696)</td>
<td>Tubulin_2 super family (E: 1.68e-07, A: cl16413)</td>
</tr>
<tr>
<td>BKP42_54900</td>
<td>Aliphatic amidase</td>
<td><em>Rhodococcus</em> sp. NJ530 plasmid unnamed 4 (Q: 100%, E: 0.0, I: 99%, A: CP034156)</td>
<td>Aliphatic amidase [<em>Rhodococcus erythropolis</em>](Q: 100%, E: 0.0, I: 100%, A: WP_095972698)</td>
<td>Acylamide amidohydrolase (E: 0.0, A: PRK13286)</td>
</tr>
</tbody>
</table>
Figure 2: Expression profile of *R. erythropolis* ACN1 under acetonitrile treatment. BKP42_54860: tubulin-like protein CetZ; BKP42_54860: hypothetical protein; BKP42_54870: tubulin/FtsZ family; GTPase domain; BKP42_54900: aliphatic amidase. Arrows: position of genes; red: forward strand; blue: reverse strand.

The expression profile of the upregulated genes revealed two independently transcribed regions (Figure 2). The first contained the amidase, while the second was an operon consisting of the three remaining genes. Between both regions, only a gene encoding for an aliphatic amidase expression-regulating protein was identified on the same strand. To exclude erroneous open reading frame (ORF) prediction during previous genome annotation, NCBI ORFfinder was used to search for all possible ORFs of the respective region (Figure S1). Combination of the obtained results and the expression profile revealed the original prediction as most plausible and strengthened assumptions on identification of a novel nitrile degrading enzyme.

**Metatranscriptomic screening approach.** Due to the distinct results obtained for *R. erythropolis* ACN1, setup was repeated with a complex microbial community. For that purpose, a compost sample was enriched in minimal medium containing either acetonitrile (ACN) or acetic acid and ammonia (ACA) as carbon and nitrogen sources. Subsequently, fresh medium with ACN or ACA was inoculated with the precultures. Obtained growth curves showed major differences between both inoculi (Figure 3). Cultures derived from the ACN inoculum and subsequently treated with ACA reached a final optical density of 1.4, whereas cultures containing ACN reached an OD₆₀₀ of 1.1. Furthermore, doubling time of the ACN-treated communities was lower than of the ACA-treated cultures. Cultures derived from the ACA inoculum behaved differently. While subsequent ACA-treatment led to fast growth and an endpoint OD₆₀₀ of 1.8, cultures treated with ACN showed a prolonged lag phase, slow growth, and reached only an optical density of 0.7.
Results

Figure 3: Growth of microbial communities under different treatments. ACA: acetic acid and ammonia; ACN: acetonitrile.

Metagenomes of both precultures as well as of all four enrichments were sequenced. After assembly, approx. 97% of the quality-filtered reads mapped to the corresponding metagenomes (Table 3) and confirmed high quality of the assembly. In addition, a length weighted median (N50) was calculated and exhibited high values for samples derived from the ACN preculture when compared to ACA-derived metagenomes.

Taxonomic analysis of the cultures revealed specific communities for each treatment (Figure 4, Data Set S2). Dominant genera in the ACN preculture were *Cupriavidus* (34.0% abundance) and *Pseudomonas* (48.8%). Similar relative abundances were recorded after subsequent ACA treatment (23.2 and 68.1%, respectively), but differed in the ACN-treated culture (74.4 and 10.1%, respectively). Community composition of cultures derived from the ACA inoculum differed significantly. The ACA inoculum mostly consisted of *Acinetobacter* (53.2%) and *Pseudomonas* (42.6%). The relative abundance of *Acinetobacter* increased approximately two-fold during further ACA enrichment (90.6%), whereas *Pseudomonas* comprises 65.8% of the subsequently ACN-enriched community. In addition, only traces of *Acinetobacter* were detected in the ACN-treated cultures. Furthermore, other organisms such as *Achromobacter*, *Cupriavidus*, and *Variovorax* exhibited a relative abundance over 1%, although they were barely present in the preculture.
Table 3: Metagenome data. Coverage based on bowtie2. Mapping reads: Percentage of quality-filtered reads mapping to the assembled metagenome.

<table>
<thead>
<tr>
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<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>ACA preculture</td>
<td>96.3 k</td>
<td>85.7</td>
<td>46.3</td>
<td>96.9</td>
<td>1,171</td>
</tr>
<tr>
<td>ACA preculture further enriched with ACA</td>
<td>33.1 k</td>
<td>30.4</td>
<td>166.6</td>
<td>98.5</td>
<td>1,041</td>
</tr>
<tr>
<td>ACA preculture further enriched with ACN</td>
<td>47.6 k</td>
<td>67.6</td>
<td>80.3</td>
<td>97.3</td>
<td>3,099</td>
</tr>
<tr>
<td>ACN preculture</td>
<td>80.5 k</td>
<td>103.8</td>
<td>51.7</td>
<td>97.2</td>
<td>2,606</td>
</tr>
<tr>
<td>ACN preculture further enriched with ACA</td>
<td>42.6 k</td>
<td>67.4</td>
<td>61.0</td>
<td>97.2</td>
<td>5,629</td>
</tr>
<tr>
<td>ACN preculture further enriched with ACN</td>
<td>56.8 k</td>
<td>75.5</td>
<td>144.9</td>
<td>97.3</td>
<td>4,325</td>
</tr>
</tbody>
</table>

Figure 4: Composition of complex communities. Community composition based on annotation of metagenomic contigs. ACA: acetic acid and ammonia; ACN: acetonitrile. Others: Organisms with abundance ≤1%.
Due to the differences in community composition between cultures derived from the ACA inoculum, transcriptome normalization was not possible. Subsequent analysis was therefore performed only for ACN-derived cultures. For normalization, the previously assembled metagenomic contigs were binned and the seven largest bins belonging to *Pseudomonas*, *Ensifer*, *Cupriavidus*, and *Flavobacterium* were used for further analysis. Lowest number of quality-filtered metatranscriptomic reads was 46.9 million and on average 60.3% of the reads mapped to one of the bins.

Table 4: Number of significantly up- or downregulated genes in the largest metagenomic bins derived from ACN cultures. Differential expression was analyzed with DESeq2 with an FDR of 0.025 and log₂FC of ≥2 as cutoff.

<table>
<thead>
<tr>
<th>Bin</th>
<th>Genus</th>
<th>No. of upregulated genes</th>
<th>No. of downregulated genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1</td>
<td><em>Pseudomonas</em></td>
<td>154</td>
<td>54</td>
</tr>
<tr>
<td>#2</td>
<td><em>Ensifer</em></td>
<td>28</td>
<td>100</td>
</tr>
<tr>
<td>#3</td>
<td><em>Cupriavidus</em></td>
<td>27</td>
<td>29</td>
</tr>
<tr>
<td>#4</td>
<td><em>Pseudomonas</em></td>
<td>186</td>
<td>65</td>
</tr>
<tr>
<td>#5</td>
<td><em>Pseudomonas</em></td>
<td>67</td>
<td>3</td>
</tr>
<tr>
<td>#6</td>
<td><em>Flavobacterium</em></td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>#7</td>
<td><em>Pseudomonas</em></td>
<td>43</td>
<td>38</td>
</tr>
</tbody>
</table>

After transcriptome normalization for each bin, differential analysis was performed with DESeq2 with the previously established parameters. In total, 505 up- and 289 downregulated genes were identified (Table 4, Data Set S3). Two of the *Pseudomonas* bins (#1 and #4) contained more than 66% of the upregulated genes, while no differentially expressed genes were identified for *Flavobacterium*. Search for genes encoding nitrile-related enzymes revealed three hits, all belonging to bin #1. The first one is an aliphatic nitrilase (log₂FC 3.2), followed by an amidase (log₂FC 3.6) and a putative amide transporter (log₂FC 5.4) (Data Set S3). For the other species, different groups of upregulated genes were identified. In case of *Cupriavidus*, five cytochrome c-related genes were upregulated, whereas another *Pseudomonas* (bin #4) contained upregulated genes belonging to the nitrate/nitrite metabolism.

In addition to a metatranscriptome analysis with focus on differential gene expression of single organisms, the normalized read counts can be pooled and analyzed as a whole. Due to the accumulated counts, non-differential expression of a gene found in several of the monitored species can become significant or vice versa (19). DESeq2 analysis on the combined normalized read counts revealed 112 up- and 77 downregulated genes under ACN treatment. Compared with the analysis of single bins, the number of differentially expressed genes was reduced by 75.3%. Of the previously identified nitrile-related genes of bin#1, only the putative amide transporter was still significantly upregulated in the combined dataset. In addition, three upregulated cytochrome c-related genes were observed, indicating a distinct effect of nitriles on this type of enzymes.
DISCUSSION

During transcriptome analysis of *R. erythropolis* ACN1, only 23 significantly up- or downregulated genes were identified. The low number was unexpected as transcriptome studies on bacteria often identify more than 100 differentially expressed genes between two conditions (10, 20, 21). Thus, nitriles seem to have a highly specific effect on the gene expression of nitrile-degrading organisms. One of the upregulated genes encoded for an amidase. Together with nitrile hydratases, these enzymes are part of a bi-enzymatic pathway for nitrile degradation, but no upregulated nitrile hydratase was found. In contrast, an upregulated operon consisting of a tubulin-like protein, a hypothetical protein, and a GTPase domain of the tubulin/FtsZ family was identified. Due to proximity to the amidase, involvement in nitrile degradation appears likely. This assumption is further supported by reports on GTPase activity of nitrile hydratase activator proteins (22, 23). Although size of the genes does not match common nitrile hydratase subunits (24–26), it seems likely that there is similar enzymatic activity. A structurally unique nitrile hydratase of *Streptomyces rimosus* is published (27) and *R. erythropolis* ACN1 could contain an additional unknown structure for these enzymes. In the end, only subcloning, overexpression, and characterization of respective genes could answer questions regarding their function.

Major difference between transcriptomes of single organisms and metatranscriptomes of complex communities is the abundance of different organisms. In case of transcriptomes, all reads can be normalized on a single genetic backbone representing the sole organism in the culture. In contrast, abundance of members of a microbial community change under different treatments (28–30). An organism highly present under the first condition can be absent under the second, making normalization and comparison of metatranscriptomes difficult (19). Consequently, cultures derived from the acetic acid-treated inoculum could not be analyzed due to major changes in community composition.

Community composition of the acetonitrile-derived cultures was suitable for bioinformatic analysis, as most genera with an abundance >1% were found in all treatments (Figure 4). After binning of the metagenomic contigs, the seven largest bins were used for subsequent normalization. Most of the bins belonged to *Pseudomonas*, but bins belonging to *Ensifer, Cupriavidus*, and *Flavobacterium* were also identified. Discrepancies between community composition and organisms identified by binning can be explained with the high number of different species in complex communities. For subsequent analysis, only the seven largest bins (based on bps) were considered. Although e.g. *Ralstonia* shows higher abundance in the community than *Flavobacterium*, this genus may comprise several species. Therefore, contigs of these different species would result in several different and probably smaller bins than obtained by a single *Flavobacterium*. Presumably as consequence of the low abundance, no
differentially expressed genes were identified for the flavobacterial bin. Modern normalization algorithms like DESeq2 and edgeR often remove low gene read counts to suppress background noise (15, 31). Although deeper sequencing would solve this problem, increasing costs must be considered during experimental design.

Quality of metagenomes is often verified by two approaches (32, 33). The first is the percentage of quality-filtered reads mapping to the final metagenome, while the second is a size-related statistic such as N50 to compare contig lengths. All acetonitrile-derived metagenomes exhibited mapping values >97%, demonstrating the high quality of the assembly. In addition, high N50 values between 2606 and 5629 were obtained, whereas other metagenomic studies often have values below 1800 (32, 34, 35). This discrepancy arises presumably from the low diversity of our microbial communities, as high coverage of a metagenome assists during the assembly of long contigs and therefore leads to high N50 values. Additional sequencing could probably further increase their quality, but based on the mapping and N50 values, the here obtained metagenomes were suitable as backbone for metatranscriptome normalization.

A first analysis of the binned metatranscriptomes demonstrated the potential of the novel screening approach. The upregulated nitrilase and amidase identified for *Pseudomonas* (bin #1) indicate nitrile degradation by at least one of the possible pathways. It can be assumed that the putative amide transporter leads to an export of amides into the medium and therefore to cross-feeding of other community members. This assumption is further supported by another *Pseudomonas* (bin #4), as its upregulated nitrate/nitrite metabolism suggests other methods than nitrile degradation to be involved in nitrogen acquisition. Thus, separate transcriptome analysis for each bin can give insights into metabolic networks of microbial communities and probably help to optimize them for bioremediation purposes.

Additional analysis of the combined metatranscriptomic read counts highlighted differences between the two analytical methods. While the upregulated nitrilase and amidase of *Pseudomonas* (bin #1) lost statistical significance, the putative amide transporter was still identified. Probably, similar genes were expressed in other members of the community but removed during bioinformatical analysis due to low read counts. In contrast, after combination of all read counts the obtained signal was strong enough to be detected. Thus, no insights into specific responses of single community members can be obtained after pooling of read counts, but larger pattern of differential expression can be observed. In conclusion, a combination of both approaches is recommended to harness the full potential of metatranscriptomics (19).
CONCLUSION AND OUTLOOK

The here presented screening approach demonstrates the potential of differential transcriptomics. During analysis of the *R. erythropolis* ACN1 transcriptomes, only four significantly upregulated genes were identified under nitrile treatment. The experimental setup seems to result in highly specific expression pattern and can therefore assist in the identification of novel biocatalysts. One of the upregulated genes belonged to an amidase, fitting our expectations, whereas the remaining upregulated genes did not belong to known nitrile-degrading enzymes. Involvement in nitrile degradation can be assumed due to the strong effect of nitriles on their expression, but further research must be done for functional assignment.

First analysis of the metatranscriptomes highlighted differences between transcriptomes of single community members and the combined read counts. Although a nitrilase most likely involved in acetonitrile degradation was identified, additional analyses must be done to uncover all enzymes involved in nitrile assimilation. Upregulated hypothetical proteins are interesting candidates for novel nitrile-degrading enzymes and especially hits found for single bins as well as the combined metatranscriptome are of major interest. In addition, binning and metatranscriptome normalization could be improved by transition to a metagenomic backbone consisting of the pooled reads of all acetonitrile-derived.

Major limitation of the presented screening approach is the enrichment of microbial communities and therefore a loss of diversity. To minimize this effect, complex communities could be grown in bioreactors. Due to better aeration and constant nutrient supply, a higher diversity should be maintained. In addition, short-term incubation of an environmental sample with a nitrile or the respective carboxylic acid and ammonia could be applied. This approach would reduce loss of diversity to a minimum but requires a high sequencing depth to gather sufficient information for subsequent analysis.

In conclusion, the here presented method provides a first insight into effects of acetonitrile on metabolism of a complex community. Additional analysis of the metatranscriptomes and especially of upregulated putative genes and the corresponding proteins would lead to identification of additional nitrile-degrading enzymes and effects. In future, this approach might probably be routinely used for identification of novel biocatalysts as it does not depend on suitable screening strategies for different substrates and can therefore be easily adapted.
MATERIAL AND METHODS

Media. M9 medium (36) was prepared containing 6 g Na₂HPO₄, 3 g KH₂PO₄, 1 g NH₄Cl, and 0.5 g NaCl per liter. After autoclaving, 10 ml ATCC trace mineral supplement, 10 ml ATCC vitamin supplement (both LGC Standards, Teddington, UK), 10 ml glucose solution (20%), 1 ml MgSO₄·7 H₂O solution (1 M), 1 ml CaCl₂ solution (14.7 g/l), and 1 ml thiamine-HCl solution (1 mg/ml) were added. For differential transcriptomics, either 25 mM acetonitrile or 25 mM acetic acid and ammonia were used as sole carbon and nitrogen sources.

Enrichment of an environmental sample. 100 g compost (pH 7.5) of the Experimental Botanical Garden Göttingen, Germany (51°33'22.6"N 9°57'16.2"E) were solved in 500 ml ddH₂O. Subsequently, the solution was filtered with a 2.7 µm GF/D glass fiber filter (Whatman, Little Chalfont, UK) and 5 ml of it were used to inoculate 45 ml M9 with acetonitrile or acetic acid and ammonia in 300 ml baffled flasks. Incubation was performed at 25 °C and 180 rpm (Innova 44 incubator). Two prevent starvation, cultures were transferred to fresh medium after reaching approx. an OD₆₀₀ of 1.0. New cultures were inoculated with an OD₆₀₀ of 0.1 and further incubated. In total, four rounds of incubation were performed.

Growth conditions. For single transcriptome studies, a preculture of R. erythropolis ACN1 (14) was grown over night in 5 ml M9 medium. For metatranscriptome studies, a previously enriched environmental soil sample was used (see before). Precultures were washed twice with M9 containing no carbon or nitrogen sources. The samples were used to inoculate 50 ml M9 with acetonitrile, or acetic acid and ammonia in 300 ml baffled flasks to an OD₆₀₀ of 0.01. Incubation was performed at 25 °C and 180 rpm using an Innova 44 incubator (New Brunswick Scientific, Nürtingen, Germany). Growth of cultures was measured regularly and samples for transcriptome sequencing were harvested during logarithmic growth phase at an OD₆₀₀ of approx. 1.0 (R. erythropolis ACN1) or 0.4 (metagenomes). For harvesting, samples were centrifuged for 1 min at 10000 × g and supernatant was discarded. Afterwards, samples were directly frozen in liquid N₂ and subsequently stored at -80 °C. All conditions were prepared as triplicates.

DNA isolation. DNA of the enrichment cultures was isolated using the Lucigen MasterPure Complete DNA and RNA Purification kit (Lucigen, Middleton, WI, USA) according to the manufacturer’s protocol. DNA of the metatranscriptome cultures was isolated with the same kit but with a modified protocol to resemble the cell disruption of the RNA isolation procedure. Pellets were dissolved in 300 µl TC buffer containing 2 µl proteinase K. Subsequently, samples were transferred to a cell mill (Mikro-Dismembrator U, B. Braun, Melsungen, Germany) containing liquid N₂ and disrupted at 1,600 rpm for 3 min. The frozen cell powder was dissolved in 1.5 ml TC buffer containing 6 µl proteinase K and DNA
Results was isolated as recommended by Lucigen. Final DNA concentration was measured using a Qubit fluorometer with dsDNA HS assay kit as recommended by the manufacturer (Invitrogen, Carlsbad, CA, USA).

**Metagenome sequencing and analysis.** DNA of triplicate cultures was pooled in equal amounts. Paired-end sequencing (2 × 300 bp) was performed on an Illumina MiSeq using the Nextera XT DNA library prep kit and v3 chemistry as described by the manufacturer (Illumina, San Diego, CA, USA). Quality trimming of the paired-end reads was done with Trimmomatic version 0.38 (37) and validated with FastQC version 0.11.7 (38). For assembly, SPAdes version 3.12 (39) was used. All contigs >200 bp were annotated with Prokka 1.13.3 (40). Coverage was calculated using Bowtie 2 version 2.3.4.2 (41) and all contigs were checked with BLASTn (42) against the NCBI non-redundant database for taxonomical assignment.

**RNA isolation.** Cell pellets were solved in 400 µl RLT buffer from the RNeasy Mini Kit (Qiagen, Hilden, Germany) containing 4 µl β-mercaptoethanol. Subsequently, cell disruption was performed in a cell mill (Mikro-Dismembrator U, B. Braun) containing liquid N₂ for 3 min at 1,600 rpm. Cell powder was dissolved in 400 µl RLT buffer containing 4 µl β-mercaptoethanol and 1.2 ml ethanol. Subsequently, the RNA isolation was done according to the manufacturer’s protocol, but with RWT instead of RW1 buffer. Final RNA concentration was measured with a Qubit using the RNA BR assay kit (Invitrogen), whereas RNA quality was controlled with a 2100 Bioanalyzer using the RNA 6000 Nano kit (Agilent Technologies, Waldbronn, Germany).

**Transcriptome sequencing.** Remaining DNA after RNA isolation was degraded with Turbo DNase (Thermo Fisher Scientific) and subsequently purified with the RNeasy MinElute Cleanup kit using 350 µl RLT buffer with 3.5 µl β-mercaptoethanol and 675 µl ethanol. After total DNA removal, rRNA depletion was done with the Ribo-Zero rRNA removal kit for bacteria (Illumina) as recommended by the supplier. Non-directed sequencing libraries were built with the NEBNext Ultra II RNA Library Prep kit (New England BioLabs) for Illumina sequencers according to the manufacturer’s protocol. Final validation of the cDNA library was done using a Qubit with the dsDNA HS assay kit (Invitrogen) and with a 2100 Bioanalyzer using the DNA High Sensitivity kit (Agilent Technologies). Single-end sequencing was done on an Illumina HiSeq 4000 (Illumina) with 1 × 50 bp for *R. erythropolis* ACN1 transcriptomes and 1 × 100 bp for metatranscriptomes.

**(Meta-)Transcriptome analysis.** Quality trimming of the single-end reads was done with Trimmomatic version 0.38 (37) and validated with FastQC version 0.11.7 (38). Normalization of the *R. erythropolis* ACN1 transcriptomes was done with R version 3.5.1 (43) and DESeq2 package version 1.20.0 (15). For normalization of the metatranscriptomes, reads were mapped against the preculture metagenome
with bowtie2 (version 2.3.4.3) (41). The mappings were used for the coverage profiles and for the number of reads assigned to a gene for the differential expression analysis. In order to compare proteins from different organisms, the protein features were unified via the KEGG ORTHOLOGY database (KO, date: 2018-10-15) (44). All proteins with a KO identifier were stored in a DIAMOND (v0.9.22.123) database (45). Proteins identified for the preculture metagenome contigs via Prokka were KO annotated using the DIAMOND database. For annotation, a target coverage of at least 90% was required and it was possible for a protein to have multiple annotations. To obtain bins of organisms, MetaBAT (version 0.32.4) (46) was used with default parameters, but --minSamples with 6 instead of 10 (Minimum number of sample sizes for considering correlation based recruiting). For the DE analysis, the 7 largest bins (bp wise) were selected, ranging from 6.8 to 2.6 Mbp. The features for the DE-analysis were obtained by counting the number of transcriptome reads overlapping with regions of the annotated KOs for each contig and summing up the KOs from contigs with the same bin id. The reads for each bin were normalized separately with DESeq2 (15) and then merged for the metatranscriptom analysis (with no further normalization). DESeq2 results with an FDR cutoff of 0.025 were subsequently filtered by just considering genes with a logarithmic fold-change of ≥2 as differentially expressed.

For graphical analysis, trimmed reads of the *R. erythropolis* ACN1 transcriptomes obtained after acetonitrile treatment were pooled. Number of reads was reduced by random subsampling by factor 10 and remaining reads were mapped with bowtie2 against the *R. erythropolis* ACN1 genome. Visualization of the results was done with TraV (47).

For additional ORF prediction, NCBI ORFfinder was used with genetic code 11. ATG and alternative initiation codons were used and nested ORFs ignored. ORFs smaller than 300 bp were not displayed.

**Accession numbers.** Biosample accession numbers for raw data of (meta)genome and (meta)transcriptome sequencing as well as assembled (meta)genomes are the following: *R. erythropolis* ACN1, SAMN05788061; acetonitrile preculture, SAMN10107935; acetonitrile preculture further enriched with acetonitrile, SAMN10112108; acetonitrile preculture further enriched with acetic acid, SAMN10112109; acetic acid preculture, SAMN10112120; acetic acid preculture further enriched with acetonitrile, SAMN10112123; acetic acid preculture further enriched with acetic acid, SAMN10112148.

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Chapter IV

General discussion
4. General discussion

Nitriles are widespread in nature and important compounds in industry. In natural systems, nitrile synthesis should equal nitrile degradation to avoid accumulation of these toxic molecules, but industrial leakage or improper disposal lead to contamination of the environment. For reduction of nitrile pollution the use of nitrile-converting enzymes and organisms as biocatalysts or in bioremediation is a promising approach. Therefore, identification of novel biocatalysts is important to expand the toolbox for green chemistry. This task is hindered by limitations of current function- or sequence-based screening approaches like (problematic) expression in heterologous hosts or inability to identify truly novel enzymes (128, 133).

In this study, a novel approach for identification of nitrile-degrading enzymes based on reverse omics techniques was developed. This approach is based on differential analysis of (meta)transcriptomic datasets without necessity of designing a suitable screening system. For this purpose, bacterial cultures are grown either with a nitrile or the corresponding products of the nitrilase reaction, carboxylic acids and ammonia. Subsequently, (meta)transcriptomes are sequenced and analyzed. Genes upregulated in presence of the nitrile could encode e.g. nitrile-degrading enzymes, transporters, or resistance mechanisms. The thereby gathered data could finally lead to the identification of novel biocatalysts. As nitriles are often toxic, understanding of their influence on microbial growth was necessary for establishing the novel screening approach. Therefore, the effect of different nitriles and their corresponding carboxylic acids on microbial growth was determined for single strains and complex communities (Chapter 3.1). One of the nitriles showed growth-supporting effects, indicating its utilization as additional nutrient source. Isolation of respective organisms led to the identification of eight novel acetonitrile-degrading bacterial strains (Chapter 3.2). In addition, a novel high-throughput nitrilase assay was developed. Subsequent screening of 70 putative nitrilases led to the identification and characterization of a new arylacetonitrilase (Chapter 3.3). Finally, a novel (meta)transcriptomic screening approach was developed. One of the previously isolated nitrile-degrading organisms was used to establish screening conditions and the subsequent bioinformatic workflow. Afterwards, a complex microbial community was identically treated and used for the first nitrilase screening approach based on differential metatranscriptomics (Chapter 3.4).

4.1. Nitriles and microbial life

Microbial communities are key players in most nutrient cycles and important for ecosystem functioning (137–143). Consequently, their natural diversity and composition (144–147) as well as influence of factors like pH (148), soil moisture (149), humidity (150), temperature (151), or salinity
(152) has been studied in detail. In addition to these environmental properties, man-made impact due to substances such as fertilizers (153), antibiotics (154), herbicides (155), or heavy metals (156) has been determined. In contrast, influence of often toxic nitriles on microbial communities is almost unknown. Besides unintended release of industrially used nitriles, they are also actively released in form of herbicides with negative impact on microbial biomass (157). Furthermore, many important crops contain cyanogenic glycosides (Table 1). For example, approximately 250 million tons of cassava with up to 1.5 g cyanide equivalent per kilogram are produced per year (158, 159). Disposed scraps of all these crops can rot in the environment and lead to increased nitrile concentrations. In addition, carboxylic acids released by nitrile degradation may also influence local biota (160, 161).

Table 1: Agriculturally important plants containing cyanogenic glycosides.

<table>
<thead>
<tr>
<th>Occurrence</th>
<th>Common name(s)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Anacardium occidentale</strong></td>
<td>Cashew</td>
<td>(162)</td>
</tr>
<tr>
<td><strong>Bambusa spp.</strong></td>
<td>Bamboo</td>
<td>(163)</td>
</tr>
<tr>
<td><strong>Cicer arietinum</strong></td>
<td>Chickpea</td>
<td>(164)</td>
</tr>
<tr>
<td><strong>Eucalyptus spp.</strong></td>
<td>Eucalyptus</td>
<td>(165)</td>
</tr>
<tr>
<td><strong>Linum spp.</strong></td>
<td>Flax</td>
<td>(166)</td>
</tr>
<tr>
<td><strong>Malus spp.</strong></td>
<td>Apple</td>
<td>(167)</td>
</tr>
<tr>
<td><strong>Manihot esculenta</strong></td>
<td>Cassava</td>
<td>(168)</td>
</tr>
<tr>
<td><strong>Pangium edule</strong></td>
<td>Keluak</td>
<td>(169)</td>
</tr>
<tr>
<td><strong>Passiflora spp.</strong></td>
<td>Passion fruit</td>
<td>(170)</td>
</tr>
<tr>
<td><strong>Phaseolus lunatus</strong></td>
<td>Lima bean</td>
<td>(171)</td>
</tr>
<tr>
<td><strong>Prunus spp.</strong></td>
<td>Almond, apricot, cherry, peach, plum</td>
<td>(162, 172, 173)</td>
</tr>
<tr>
<td><strong>Sambucus spp.</strong></td>
<td>Elderberry</td>
<td>(173, 174)</td>
</tr>
<tr>
<td><strong>Simmondsia chinensis</strong></td>
<td>Jojoba</td>
<td>(175)</td>
</tr>
<tr>
<td><strong>Sorghum spp.</strong></td>
<td>Sorghum</td>
<td>(176)</td>
</tr>
<tr>
<td><strong>Trifolium repens</strong></td>
<td>White clover</td>
<td>(177)</td>
</tr>
<tr>
<td><strong>Vicia sativa</strong></td>
<td>Common vetch</td>
<td>(173)</td>
</tr>
</tbody>
</table>

For a first insight into possible effects of nitriles and their corresponding carboxylic acids, influence on bacterial growth was studied (Chapter 3.1). Previous toxicity experiments on mammals have shown diverging effects of structurally different nitriles (63). Consequently, structurally diverse substrates were chosen in this study. In total, nine different mononitriles, dinitriles, cyanohydrin derivatives, arylacetonitriles, aromatic nitriles, cyclohexane nitriles (Figure 1), and their corresponding carboxylic acids were selected.
After determination of nitrile toxicity on *Agrobacterium tumefaciens*, *Bacillus subtilis*, *Corynebacterium glutamicum*, and *Escherichia coli* using solid media, effect of the nitriles and carboxylic acids on a complex microbial community was analyzed. Compost was chosen as starting material to increase chance for viable communities due to its higher diversity compared to aquatic environments (178). Common methods for extraction of similar communities use serial dilutions (179) or sonication and subsequent low-speed centrifugation (180, 181). The first method leads to dilution of microorganisms and therefore loss of biomass, whereas the second method can severely damage the cells and reduce number of viable organisms (181). Consequently, a different approach was tested. For that purpose, compost was solved in H₂O and filtered with a 2.7 µm glass fiber filter to remove matrix particles. Pore size of the filter was chosen to allow flowthrough of presumably the largest bacterium found in German soil, *Bacillus megaterium* (182). 16S rRNA gene analysis of the unfiltered and filtered soil sample revealed very similar community compositions with more than 500 different organisms at genus level and identical Simpson indices of diversity (SID) values. Thus, it is indicated that the here established method is suitable for removal of compost or soil particles in future studies of similar communities.

During toxicity determination, two important results were retrieved. First is the high toxicity of 4-hydroxybenzonitrile, leading to suppressed growth and reduced diversity. Structurally similar herbicides like bromoxynil and ioxynil show similar impact on microbes (157, 183). Thus, it can be assumed that the use of 4-hydroxybenzonitrile-derived herbicides has severe influence on soil microbiota. More than 1,300 tons of bromoxynil have been used in the US in 2016 (184) and most likely the use of ioxynil is even more widespread as it is also available for private households. Transition to less toxic herbicides should be considered to increase quality of soil microbiota and therefore crop yield (185).

The second important result is the lethal effect of acetone cyanohydrin. This cyanohydrin is the degradation product of linamarin, a cyanogenic glycoside found in cassava, lima beans, and flax (186–188). Due to large-scale cultivation of these crops, release of acetone cyanohydrin by rotting...
scraps could be a source for environmental contamination and hazardous for local microbiota. Nevertheless, further studies must be performed for estimation of the impact of cyanogenic crops on the environment.

In addition to specific levels of toxicity, a higher nitrile susceptibility of Gram-negatives compared to Gram-positives was recorded. Research on tolerance or resistance of both bacterial groups to antibiotics, hydrocarbons, cold plasma, or special nanoparticles has been performed before (189–194), but this is the first time that nitrile-susceptibility has been studied. Due to the small number of tested compounds and organisms, these findings can just serve as tendency. Further studies including other nitriles and several different communities are necessary.

4.2. Acetonitrile-degrading bacteria

Acetonitrile is an industrially important nitrile often employed as solvent. It is used during the production of butadiene (195), a key solvent in pharmacy (196), and part of battery electrolytes (197). In addition, it is important for the synthesis of oligonucleotides (198) and often used for liquid chromatography (199). Nevertheless, it exhibits higher toxicity than other solvents such as methanol or acetone and is regarded as problematic waste (199, 200). Conventional treatment of acetonitrile waste can be done by addition of large quantities of sodium hydroxide (201), with supercritical water and H₂O₂ (202), by photolysis (203), or via combustion in a hazardous waste plant (201). As most of these processes need unfavorable chemicals or conditions, interest in bioremediation of acetonitrile-containing waste is increasing (204–206). During analysis of nitrile toxicity on microbial communities (Chapter 3.1), cultures containing acetonitrile exhibited higher optical density than the control. As nitrile-metabolization and therefore presence of nitrile-degrading enzymes could be assumed, isolation of respective organisms was performed (Chapter 3.2).

Eight unique strains belonging to *Flavobacterium*, *Pseudomonas*, *Rhodococcus*, and *Variovorax* were isolated. Four of the eight isolates belonged to *Pseudomonas*, demonstrating its well-known acetonitrile-degrading abilities (207–209). In addition, one *Rhodococcus* isolate was identified. This genus is probably the most prominent bacterial genus for nitrile bioremediation and biocatalysis. It can degrade a large number of compounds including acetonitrile (210–212), also reflecting by the increasing number of related patents (213). Despite these common nitrile degraders, one isolate of *Variovorax* and two isolates of *Flavobacterium* were obtained. Nitrile-degrading ability of *Variovorax* was described before (214, 215), but only weak hydrolysis of nicotinonitrile by *Flavobacterium* has previously been observed (216). Due to missing genomes or enzymatic studies, the pathway involved in nicotinonitrile degradation remains unknown. Although no nitrilases or nitrile hydratases could be annotated in the genomes of the flavobacterial isolates, these results indicate that nitrile degradation...
is even more widespread than previously known. Further studies are required to unravel the underlying mechanism and increase the number of nitrile-degrading biocatalysts.

4.3. High-throughput nitrilase assays

In addition to identification of nitrile-degrading bacteria (Chapter 3.2), characterization of putative nitrile-degrading enzymes is of increasing importance. To analyze the growing amount of putative nitrilases and nitrile hydratases found in databases or hiding in metagenomic libraries, simple and efficient high-throughput assays are necessary. For that purpose, several assays based on three different principles have been developed (217). The first group uses indicators like bromothymol blue, bromocresol purple, or phenol red to measure a pH shift caused by the carboxylic acid or the ammonia released during nitrile degradation (122, 218, 219). These methods are easy to perform and the respective color shifts can be detected without additional equipment. However, their success depends on the strength of the released acid. In case of acids equally strong as the basic ammonia, no pH shift will occur and therefore prevent the detection of nitrile-degrading enzymes. The second group of high-throughput assays focuses on the released carboxylic acids (220, 221). Due to the structural diversity of carboxylic acids, these methods are often limited to specific molecules, making the screening of broad substrate libraries impossible. The third group of assays detects the ammonia released by nitrilase reactions. Besides screenings based on the established Berthelot reaction (217), a fast and simple assay using CoCl₂ as indicator is published (222). Drawback of the first is the use of highly toxic phenol hypochlorite mixtures, whereas the second is not sensitive enough for detection of low amounts of ammonia.

To assist functional screening of large amounts of putative nitrile-degrading enzymes, a novel nitrilase high-throughput assays was developed during the here presented work (Chapter 3.3). It is based on well-established NAD(P)H-coupled systems, which monitor the conversion of NAD(P)H to NAD(P)⁺ or vice versa (223–226). In case of nitrilases, the released ammonia is converted by a glutamate dehydrogenase (GDH) to glutamate (Figure 2). This reaction involves the oxidation of NADH to NAD⁺ and is used to determine nitrile degradation.

![Nitrilase Reaction](image.png)

**Figure 2: Reactions of a NADH-coupled nitrilase assay.**
In contrast to a previously published NADH-coupled nitrilase assay (132), the novel approach does not depend on long incubation times. For that purpose, nitrile degradation and NADH conversion were coupled in a single mixture. By using a plate reader, real-time conversion of the cofactor is monitored in 96-well plates, providing direct insights into nitrilase efficiency for different substrates. Furthermore, the assay was performed with crude extract containing the overexpressed putative enzymes, allowing the simple and fast screening of large (meta)genomic libraries.

Unspecific NADH conversion was observed during first tests of the assay with an already characterized nitrilase. Most likely, unintended NADH oxidation by other proteins of the crude extract caused NAD\(^+\) formation. For example, members of the tricarboxylic acid (TCA) cycle like malate dehydrogenases can also convert NADH to NAD\(^+\) and therefore influence the obtained signal (227). Consequently, a sharp threshold is recommended to reduce number of false positives. In addition, every reaction exhibited a turning point after which the optical density was increasing. This observation was independent of the used substrate and most likely caused by the reverse reaction of NAD\(^+\) to NADH. In consequence, this assay can lead to wrong conclusions when performed with single measurements. It is not possible to differentiate between nitrilase activity, side reactions of the crude extract, and regeneration of the cofactor for a single data point. Only the full reaction curve obtained by continuous measurements with a fast plate reader allows assessments of nitrile degradation.

Overall, the here presented workflow allows the simple and fast verification of putative nitrilases identified during sequence-based screenings. Optimization of the bioinformatic analyses could further improve the outcome of nitrilase screenings and advance identification of novel biocatalysts. During \textit{in silico} analyses of the 70 putative nitrilases used as input, almost 50\% of the enzymes were excluded from further analysis due to their low quality. Still, only putative enzymes clustering with characterized nitrilases obtained from SWISS-Prot showed enzymatic activity. Therefore, it can be recommended to discard all “unique” putative enzymes when time restrictions apply or the number of putative nitrilases is too large for handling. Nonetheless, it has to be considered that this additional filtering will also reduce the chance for identification of truly novel enzymes. Furthermore, prediction of substrate specificity based on sequence homology could help to identify possible genes of interest before wet lab work even starts. Such an approach was already published with focus on dinitriles and led to promising results (228). In combination with the novel high-throughput screening, identification of nitrilases of interest could be significantly improved and support the increasing industrial demand for these biocatalysts.
4.4. Characterization of a novel arylacetonitrilase

Although nitrilases are an important class of enzymes for various industrial processes, only a small number has been characterized. Curated enzyme databases like SWISS-Prot contain thousands of phosphatases, esterases, or lipases, but less than 100 nitrilases. Relationship between sequence, structure, and function is nearly unknown because of the low number of characterized nitrilases (75). Consequently, one of the nitrilases identified with the novel high-throughput assay was characterized in detail to support knowledge and understanding of these enzymes. Due to its unique substrate specificity, the phenylacetonitrile-degrading nitrilase was chosen for this task (Chapter 3.3).

**Table 3: Properties of characterized bacterial arylacetonitrilases.** Mass refers to mass of homomers; activity refers to specific activity for phenylacetonitrile.

<table>
<thead>
<tr>
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<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Alcaligenes faecalis</em> JM3</td>
<td>7.5</td>
<td>45</td>
<td>44</td>
<td>85.0</td>
<td>(229)</td>
</tr>
<tr>
<td><em>Alcaligenes faecalis</em> ATCC 8750</td>
<td>7.5</td>
<td>40-45</td>
<td>32</td>
<td>30.0</td>
<td>(230)</td>
</tr>
<tr>
<td><em>Alcaligenes faecalis</em> ZJUTB10</td>
<td>7.5</td>
<td>40</td>
<td>44</td>
<td>41.9</td>
<td>(231)</td>
</tr>
<tr>
<td><em>Alcaligenes</em> sp. MTCC 10675</td>
<td>6.5</td>
<td>50</td>
<td>60</td>
<td>65</td>
<td>(232)</td>
</tr>
<tr>
<td><em>Bradyrhizobium japonicum</em> USDA110</td>
<td>-</td>
<td>-</td>
<td>37</td>
<td>5.3</td>
<td>(233)</td>
</tr>
<tr>
<td><em>Luminiphilus syltensis</em> NORS1-1B</td>
<td>7.0</td>
<td>40</td>
<td>43</td>
<td>89.2</td>
<td>(234)</td>
</tr>
<tr>
<td><em>Pseudomonas fluorescens</em> EBC191</td>
<td>6.0-6.5</td>
<td>50-55</td>
<td>38</td>
<td>67.7</td>
<td>(235)</td>
</tr>
<tr>
<td><em>Pseudomonas putida</em> MTCC 5110</td>
<td>7.0</td>
<td>40</td>
<td>43</td>
<td>11.2</td>
<td>(78)</td>
</tr>
<tr>
<td>Compost metagenome</td>
<td>6.0</td>
<td>50</td>
<td>36</td>
<td>13.9</td>
<td>This study</td>
</tr>
</tbody>
</table>

Based on its substrate specificity, the novel nitrilase belongs to the class of arylacetonitrilases (EC 3.5.5.5). Arylacetonitrilases are probably the industrially most important group of nitrile-degrading enzymes as they allow the conversion of mandelonitrile to mandelic acid (110). With less than 10 respective entries in databases like SWISS-Prot, they are the smallest subgroup of nitrilases and to the best of my knowledge, only *Alcaligenes*, *Bradyrhizobium*, *Luminiphilus*, and *Pseudomonas* contain characterized bacterial arylacetonitrilases (Table 3). Consequently, several studies with focus on understanding and optimization of these enzymes have been performed (236–241). Nevertheless, identification of novel arylacetonitrilases will provide new opportunities and insights into sequence-function relationships that cannot be easily obtained by other experiments. For example, based on protein size as well as pH and temperature optimum, the here characterized enzyme is closely related to the arylacetonitrilases of *B. japonicum* USDA110 and *P. fluorescens* EBC191 (Table 4). These results are further supported by sequence similarity, as the respective enzymes build a separate branch of arylacetonitrilases (Figure 4). In contrast, the overall substrate specificity of the novel arylacetonitrilase is similar to that of *P. putida* MTCC 5110, although their sequences are not closely related.
Figure 4: Phylogenetic tree containing all identified novel nitrilases. 500 bootstrap replicates were calculated and the tree was condensed at values below 50%. No nitrilase sequence was available for *Luminiphilus syltensis* NOR5-1B. WAG algorithm including gamma sites was chosen for calculation of the phylogenetic tree (242). For general parameters and references for *R. rhodochrous* M8 amidase, see material and methods section of chapter 3.1. NCBI accession number for protein sequences are in brackets, for references see Table 3.

Besides its unusual sequence-function relationship, the novel nitrilase exhibits other non-common features like optimum activity at high temperatures and over a broad pH range. Furthermore, it exhibits a unique long-term stability as more than 80% of initial activity remained after two weeks of storage. In contrast, arylacetonitrilases of *A. faecalis* JM3 and *P. fluorescens* EBC191 showed only 40% and 25% remaining activity after one or two weeks under similar conditions, respectively (229, 235). In conclusion, the here identified and characterized nitrilase leads to a better understanding of sequence-function relationship of these rare and valuable enzymes and is an interesting candidate for industrial applications.

4.5. A novel (meta)transcriptomic screening approach

Screening for novel enzymes is nowadays often done by function- or sequence-based metagenomic approaches (see Chapter 2.7). Both have their specific drawbacks such as problems due to expression in heterologous hosts or inability to identify truly novel enzymes. In addition, they cannot identify complex networks sometimes involved in metabolization of a substrate (243). Differential transcriptomic approaches are without these inherent limitations. Microbial cultures are grown under different conditions and their transcriptomes are isolated and sequenced. Comparison of the expression profiles highlights genes up- and downregulated for each treatment, leading to identification of genes encoding enzymes or pathways specific for the tested conditions. Although this approach seems to be promising for identification of novel biocatalysts, most research focuses on clinical aspects like human diseases or different pathogens (244–252). In contrast, only a few studies with focus on industrial topics such as increased solvent tolerance of living cells or identification of novel biocatalysts have been performed (136, 253–255).
Important for differential transcriptomics is proper normalization of the transcriptomes. Without these steps during data analysis, obtained results could be misleading or wrong. Consequently, different approaches for normalization of transcriptomic data have been developed and were extensively reviewed (256–259). Several of these methods depend on count-based normalization, but compensate only for differences in sequencing depth (256). When comparing two conditions with equal strength in gene expression but double the number of expressed genes in the first condition, these approaches will introduce a severe bias. After total count normalization, only half the number of reads per gene will be assigned in the first treatment, distorting the outcome of the analysis (260). To account for this problem, new statistical models have been proposed. They assume that most genes between two samples are not differentially expressed and calculate a correction factor for all read counts to fulfill this hypothesis (256). Most prominent implementations of this approach can be found in edgeR (261) and DESeq2 (262) and both methods are nowadays recommended for transcriptome analysis (256, 257).

When trying to apply standard transcriptome normalization methods like DESeq2 to metatranscriptomes, a major problem occurs. During work with a single strain, abundance of this organism in the culture is always 100%. In contrast, abundance of organisms can significantly change in complex communities between two treatments (263–265), as also found in the here presented study. If an organism would be absent during first treatment and present during second, all its genes would appear upregulated under the second treatment during subsequent analysis. To prevent this community-based bias, an additional normalization step is necessary (266). For that purpose, the metagenomic backbone must be divided into bins consisting of single species. Subsequently, transcriptomic reads can be assigned to their biological origin and are individually normalized for each bin. Afterwards, the normalized reads can be combined to the final metatranscriptomic dataset and analyzed with algorithms like DESeq2. To the best of my knowledge, the here presented metatranscriptomic study is the first using this thorough normalization approach. Previous studies dealing with differential metatranscriptomics did not normalize their data based on community composition and may therefore contain erroneous results or assumptions (266–273).

To establish the novel transcriptome-based screening approach, acetonitrile was used as substrate of choice (Chapter 3.4). During previous toxicity analysis (Chapter 3.1), no growth-suppression was observed even at 25 mM concentration. Furthermore, cultures treated with acetonitrile exhibited high diversity, indicating no toxic effect on many microorganisms. In conclusion, unintended expression of stress response genes should be minimal for this compound when compared to other tested nitriles. For a proof of principle, the previously isolated acetonitrile-degrading *Rhodococcus erythropolis* ACN1 (Chapter 3.2) was used. When comparing the transcriptomes of *R. erythropolis* ACN1 after treatment with acetonitrile or acetic acid and ammonia, 23 differentially expressed genes were identified.
Apparently, acetonitrile has a minor effect on general gene expression and is therefore a promising substrate for metatranscriptome-based nitrilase screenings. In addition, only four of the differentially expressed genes were upregulated in the acetonitrile-containing sample. Most likely all of them are involved in nitrile degradation, demonstrating the potential of the tested setup.

Analysis of the metatranscriptomes revealed many differentially expressed genes. Comparing the *R. erythropolis* ACN1 transcriptome with the metatranscriptome, number of differentially expressed genes per species increased from 23 (4 up- and 19 downregulated) to 113 (72 up- and 41 downregulated). Presumably, cross-feeding and metabolic networks across different community members contribute to these differences (274–277). Although these effects could hinder the fast identification of novel biocatalysts, they allow insights into microbial interaction. In addition, the here presented approach may increase knowledge on resistance mechanisms. An example can be seen for *Cupriavidus* (species #3) with several upregulated cytochrome c-related genes. Nitrile-induced inhibition of cytochrome c oxidases has been proposed for eukaryotes (Chapter 2.3), but mode of action in prokaryotes is unknown. The metatranscriptomic data solidify assumptions on similar mechanisms in bacteria as overexpression of these essential genes may help to compensate the inhibition. In addition to putative resistance mechanisms, a highly upregulated nitrilase-encoding gene was identified. It can be assumed that this enzyme acts on acetonitrile, making it the first nitrile-degrading biocatalyst discovered by a transcriptomic approach. Nevertheless, further characterization of the respective enzyme is necessary to confirm this assumption. For that purpose, the previously established high-throughput assay can be used (Chapter 3.3), as it would allow the fast and simple screening of the putative nitrilase with several substrates.

In conclusion, the here presented screening demonstrates the potential of differential metatranscriptomics. Although first analysis revealed just two enzymes involved in nitrile degradation, valuable insights into cross-feeding and putative resistance mechanisms were obtained. In addition, analysis of upregulated hypothetical proteins and screening with a high-throughput nitrilase assay could lead to identification of novel biocatalysts. Screening of nitrilases was challenging due to toxicity of the substrate, but should be simple for other compounds like lipids, alcohols, or esters. Therefore, this approach might lead to identification of several novel classes of biocatalysts and is a promising method to support the demands and goals of green chemistry.
Chapter V

General references
5. **General references**


6. Appendix

6.1. Publications


*Shared first author*

6.2. Posters at conferences


6.3. Declaration of independent work

I, Richard Egelkamp, hereby declare that the doctoral thesis

“Novel approach for identification of biocatalysts by reverse omics techniques”

has been written independently and with no other sources and aids than quoted. Moreover, I have not used unauthorized assistance and have not submitted this thesis previously in any form for another degree at any institution or university.

Richard Egelkamp

Göttingen, 21.12.2018
6.4. Acknowledgment

Every research project has its ups and downs and a PhD thesis can feel like an emotional roller coaster. On good days, the Nobel Prize seems within reach, while on bad days, the project feels like an impossible task. This thesis would not have been written without the continuous support and believe of many people and I wanted to save the last page for them.

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