Isotope fractionation pinpoints membrane permeability as barrier to atrazine biodegradation in Gram-negative *Polaromonas sp. Nea-C*

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ABSTRACT: Biodegradation of persistent pesticides like atrazine often stalls at low concentrations in the environment. While mass transfer does not limit atrazine degradation by the Gram-positive *Arthrobacter aurescens* TC1 at high concentrations (> 1 mg∙L⁻¹), bioavailability limitations have recently been observed at trace concentrations (< 0.1 mg∙L⁻¹). To assess the bioavailability constraints on biodegradation, the roles of cell wall physiology and transporters remain imperfectly understood. Here, compound-specific isotope analysis (CSIA) demonstrates that cell wall physiology (i.e. the difference between Gram-negative and Gram-positive bacteria) imposes mass transfer limitations in atrazine biodegradation even at high concentrations. Atrazine biodegradation by Gram-negative *Polaromonas sp.* Nea-C caused significantly less isotope fractionation (ε(C) = -3.5 ‰) than expected for hydrolysis by the enzyme TrzN (ε(C) = -5.0 ‰) and observed in Gram-positive *Arthrobacter aurescens* TC1 (ε(C) = -5.4 ‰). Isotope fractionation was recovered in cell free extracts (ε(C) = -5.3 ‰) where no cell envelope restricted pollutant uptake. When active transport was inhibited with cyanide, atrazine degradation rates remained constant demonstrating that atrazine mass transfer across the cell envelope does not depend on active transport but is a consequence of passive cell wall permeation. Taken together, our results identify the cell envelope of the Gram-negative bacterium *Polaromonas sp.* Nea-C as relevant barrier for atrazine biodegradation.
Introduction

Groundwater contamination by micropollutants is a prominent challenge of our time. Since ground and surface waters represent an important drinking water resource, the presence of micropollutants is of concern not only for ecosystems, but also for human health. Because of their ubiquitous release and their low concentrations, however, evaluating the fate of micropollutants in the environment is complex. Pharmaceuticals are discharged into the environment with wastewater treatment effluents and pesticides used in agriculture even directly leach into groundwater on a large scale. EU regulations on drinking water quality set a maximum concentration of 0.1 μg/L for pesticides and their degradation products (European Union Drinking Water Directive, 98/83/EC). How difficult it is, however, to relate successful biodegradation in the lab to the fate of pesticides in the environment, is illustrated by the herbicide atrazine: even though banned in the EU as long ago as 2003, atrazine and its metabolites are still the groundwater contaminants most frequently detected above this threshold concentration. The underlying bottlenecks of biodegradation at trace concentrations which cause this persistence have eluded researchers for years. Even though atrazine is initially adsorbed and retained to some extent on soil and sediments, the pesticide becomes available at low concentrations (μg/L) for atrazine degrading bacteria. For such a situation, competing models claim that it is either mass transfer (uptake into microbial cells) which puts a limit to otherwise rapid enzymatic transformation, or that physiological limitations (enzyme activity, downregulation) prevail. Compound-specific isotope analysis (CSIA) provides a way to directly visualize the rate-determining step of pollutant biodegradation: Chemical bond breakage during pollutant degradation is slower when the bond contains a heavy isotope since the respective activation energy is higher. Therefore, the remaining pollutant molecules contain on average increasingly more heavy isotopes as an enzymatic
reaction proceeds.\textsuperscript{15} This trend can be described by relating the change in isotope ratios $(R_t/R_0)$ to the fraction of the remaining pollutant $f$ according to the Rayleigh equation\textsuperscript{16, 17} (1)

$$\ln \left( \frac{R_t}{R_0} \right) = \varepsilon \times \ln(f)$$

where the enrichment factor $\varepsilon$ reflects the incremental isotope fractionation during transformation. This isotope effect, however, can only be observed if substrate molecules experiencing the isotopic discrimination during the enzymatic reaction in the cytosol diffuse back out into the bulk solution, where the isotope ratio is assessed. Thus, any partially rate-determining step preceding the irreversible enzymatic turnover (e.g. mass transfer) will lead to a reduced exchange of “enriched” substrate molecules to the outside of the cell. As a consequence, the observable isotope enrichment factor $\varepsilon$ will be smaller, since the reduced exchange masks the isotope fractionation of the enzymatic reaction.\textsuperscript{18, 19} Masked isotope fractionation due to mass transfer limitations is well understood from photosynthesis,\textsuperscript{20, 21} sulfate reduction,\textsuperscript{22-24} or nitrate reduction.\textsuperscript{25, 26} The same effect has previously been demonstrated for organic pollutants taken up by active transport\textsuperscript{27} and for non-polar chlorinated ethenes.\textsuperscript{19, 28} A conceptual framework has been brought forward by Thullner \textit{et al.} to mathematically predict the effect for passive permeation of organic pollutants through a biological double membrane.\textsuperscript{29, 30} Based on these studies, we recently discovered that cell wall permeation was not relevant for atrazine biodegradation by \textit{Arthrobacter aurescens} TC1 at high concentrations, but became suddenly rate-limiting at low concentrations (low µg/L range).\textsuperscript{31} This finding is challenged by earlier observations by Meyer \textit{et al.} that even at high concentrations, isotope fractionation in atrazine degradation varied significantly between bacterial strains catalyzing the same reaction.\textsuperscript{32} Usually, the isotope fractionation factor is assumed to be characteristic for a specific transformation pathway if the underlying enzyme reaction is identical.\textsuperscript{33-35} A compelling clue to explain the results of Meyer \textit{et al.}, is, therefore,
the fact that differences exist between Gram-negative and Gram-positive bacterial strains. The additional outer membrane in Gram-negative strains possibly constitutes an additional barrier for mass transfer which can mask the enzymatic isotope fractionation. Indeed, Renpenning et al. observed that carbon isotope fractionation during chlorinated ethene degradation differed for Gram-positive and Gram-negative bacteria and depended on the integrity of the cell envelope. In the case of Meyer et al., however, this proposed causal relationship could not be uniquely pinpointed because different enzymes of the same family (AtzA vs. TrzN) were involved. Consequently, it could not be excluded that the observed variability may, alternatively, be attributable to subtle variations in transition state structures.

We, therefore, systematically addressed the question in our study by exploiting the opportunity that the Gram-negative bacterium, *Polaromonas sp.* Nea-C, harbors the same set of intracellular atrazine degrading enzymes as the Gram-positive *A. aurescens* TC1 (TrzN, AtzB, AtzC). Atrazine hydrolysis by TrzN ($K_M = 19 \mu M, k_{cat} = 5.5 s^{-1}$) proceeds via initial protonation of the ring nitrogen and subsequent hydrolysis of the C-Cl bond. Further, enrichment factors of the degradation reaction in whole cells are similar to those of the degradation with purified TrzN without cell envelope meaning that atrazine degradation by whole cells of Gram-positive *A. aurescens* TC1 is not mass transfer limited. We compared the isotope fractionation during atrazine degradation with intact cells of Gram-negative *Polaromonas sp.* Nea-C – a scenario in which mass transfer across the cell envelope can matter – relative to degradation with Gram-positive *A. aurescens* TC1, cell free extracts of Gram-negative *Polaromonas sp.* Nea-C or purified TrzN enzyme, three scenarios in which mass transfer is absent. Furthermore, we addressed the possibility of active transport to clarify whether passive membrane permeation of atrazine is sufficient to provide enough influx for those bacteria to sustain growth. To this end, we investigated whether atrazine
degradation rates of *Polaromonas sp.* Nea-C and *A. aurescens* TC1 were affected when active transport was inhibited by the respiratory chain inhibitor potassium cyanide (KCN).

**Experimental section**

**Chemicals**

A list of chemicals used can be found in the supporting information.

**Cultivation of bacteria**

*Polaromonas sp.* Nea-C was kindly provided by Fabrice Martin-Laurent (Microbiologie du Sol et de l’Environnement, INRA, France) and *Arthrobacter aurescens* TC1 was kindly provided by Larry Wackett (The BioTechnology Institute, University of Minnesota, USA).

All strains were grown in liquid mineral salt medium (MSM) containing a nitrogen source (composition see supporting information). Excess atrazine above the solubility limit was added in solid form to a concentration of 500 mg/L to provide enough nutrient for high cell densities. Cultures were incubated at room temperature (25°C).

**Atrazine degradation with whole cells of *Polaromonas sp.* Nea-C for isotope analysis.**

Growth of a freshly inoculated culture (500 mL) of *Polaromonas sp.* Nea-C was followed by monitoring the optical density (OD$_{600}$). During exponential phase (OD$_{600} = 0.05$) cells were pelleted by centrifugation (Heraeus Megafuge 40R, Thermo Scientific, TX-1000 rotor, 3700 g, 30 min, 4°C) and washed twice in 50 mL MSM to remove the remaining atrazine. After those washing steps, the cell pellet was resuspended in 500 mL of fresh MSM containing 30 mg/L atrazine. The atrazine concentration was close to the solubility limit of 33 mg/L (see media preparation in the SI) to maximize the amount of substance per volume and, hence, to minimize the necessary sample volume for reliable isotope analysis (see below). The degradation experiment lasted approximately 24 h and the atrazine concentration was monitored by HPLC-UV (see below). For each of the three biological replicates, 5 samples for isotope analysis were taken (20 mL in the beginning and 50, 70, 150, 200 mL at...
approximately 50 %, 75 %, 85 %, and 95 % atrazine consumption respectively). The degradation reaction was stopped by sterile filtration with a regenerated cellulose membrane filter (pore size 0.2 µm, diameter 47 mm; GE Healthcare ltd., UK). The biomass and the filter volume were not extracted, as their volume (< 0.5 mL) is negligible compared to the filtrate (> 20 mL). The filtrate was extracted three times with 10 % (v/v) dichloromethane. The combined dichloromethane extracts were evaporated under an air stream and the samples were reconstituted in 100 µL ethyl acetate for GC-IRMS measurements (see below).

**Preparation of cell free extracts of *Polaromonas sp. Nea-C***.

*Polaromonas sp. Nea-C* cells were grown and harvested as described above. The cell pellet was resuspended in 5 mL of fresh MSM and put on ice. Cell membranes were disrupted in two passages by a French pressure cell (American Instrument Company, USA, 3/8” piston diameter, 20000 Psi). Remaining whole cells and cell fragments were removed by sterile filtration with a regenerated cellulose membrane filter (pore size 0.2 µm, diameter 47 mm; GE Healthcare ltd., UK) and the extract was stored on ice for a short time for the degradation experiment.

**Atrazine degradation with cell free extracts of *Polaromonas sp. Nea-C* for isotope analysis.**

The 5 mL concentrated cell free extract was diluted in 250 mL of fresh MSM containing 30 mg/L atrazine. The atrazine concentration over time was monitored by HPLC-UV (see below) by taking samples for 4 h. For each of the three biological replicates, 5 samples for isotope analysis were taken (10 mL in the beginning and 15, 35, 60, 120 mL at approximately 60 %, 80 %, 90 %, and 95 % atrazine consumption respectively) and the degradation reaction was stopped by extracting atrazine with three times 10 % (v/v) dichloromethane. The extracts were concentrated for GC-IRMS as described above.

**Atrazine degradation rates with and without respiratory chain inhibitor KCN.**
Growth of freshly inoculated cultures (50 mL) of both, *Polaromonas sp. Nea-C* and *A. aurescens* TC1, was followed by the OD$_{600}$ and cell numbers per mL were estimated for both strains with $8 \cdot 10^8$ cells∙mL$^{-1}$∙OD$_{600}$. The cells were harvested as described above and the cell pellet was resuspended in 50 mL of fresh MSM. For both species, each of the three biological replicates was split in 2 x 25 mL cell suspensions to get the same biomass for the inhibited and the non-inhibited degradation experiment. To inhibit the respiratory chain, 0.25 mM KCN was added to one cell suspension. Afterwards, atrazine was added to both cell suspensions at a concentration of 3 mg/L. A small initial atrazine concentration was chosen to ensure short degradation times to rule out growth of the non-inhibited cells during the experiment. The atrazine concentration over time was monitored by HPLC-UV for 4 h (see below). Because *Polaromonas sp. Nea-C* and *A. aurescens* TC1 might have different TrzN abundances, we only compared each strain with and without inhibition and not the degradation kinetics of *Polaromonas sp. Nea-C* versus *A. aurescens* TC1.

**Determination of the atrazine concentration by HPLC-UV.**

Atrazine concentrations were measured using a Prominence HPLC system (Shimadzu Corp., Japan) together with a 100 x 4.6 mm Kinetex 5 μ Biphenyl 100 Å column equipped with a SecurityGuard ULTRA Biphenyl cartridge (both Phenomenex Inc., USA). The injected sample volume was 10 μL. Peak separation was achieved by 1 mL/min isocratic flow of a mixture of 51 % 5 mM KH$_2$PO$_4$ buffer at pH 7 and 49 % methanol, respectively, for 9 min. The compounds were detected by UV absorbance at 222 nm and the peaks were quantified using LabSolutions V 5.71 SP2 (Shimadzu Corp., Japan). External calibration was conducted with atrazine dissolved in 25 % methanol and 75 % water in the following concentrations: 0.5, 4, 12, 35 μg/L.

**Carbon and nitrogen isotope measurements with GC-IRMS**
The method was adapted from Reinnicke et al.\textsuperscript{39} The GC-IRMS system consisted of a TRACE GC Ultra gas chromatograph (GC; Thermo Fisher Scientific, Milan, Italy) linked to a Finnigan MAT 253 isotope ratio mass spectrometer (IRMS) (Thermo Fisher Scientific, Germany) by a Finnigan GC Combustion III Interface (Thermo Fisher Scientific, Germany). Helium (grade 5.0) was used as carrier gas and the split injector was kept at 250°C with a 1:10 split at a flow rate of 1.4 mL/min. The samples were injected using a GC Pal autosampler (CTC, Switzerland) onto a 60-m DB-5 (30 m × 0.25 mm; 1 μm film; Restek GmbH, Germany) analytical column. Isotope values were determined as δ\textsuperscript{13}C and δ\textsuperscript{15}N values in per mill relative to Vienna PeeDee Belemnite (VPDB)\textsuperscript{40} and Air-N\textsubscript{2}\textsuperscript{41} The δ\textsuperscript{13}C and δ\textsuperscript{15}N values were assessed in relation to a monitoring gas (CO\textsubscript{2} and N\textsubscript{2}, respectively) which was measured alongside each run at the beginning and the end. Calibration of monitoring gases was performed in a Finnigan MAT Delta S isotope ratio mass spectrometer with dual inlet system (Thermo Fisher Scientific, Germany). The gases were measured against VPDB and air, respectively, by use of international reference materials: the CO\textsubscript{2} gases RM 8562, RM 8563, and RM 8564 for CO\textsubscript{2} and NSVEC (N\textsubscript{2} gas) for N\textsubscript{2}. Reference standards were provided by the IAEA. The GC oven started at 65 °C (hold 3 min), ramp 25 °C/min to 190 °C This was followed by a temperature ramp of 15 °C/min to 270 °C which was kept for 20 min.

\textbf{Modelling of the isotope fractionation during the degradation.}

In the absence of the cell envelope, the bioavailable concentration is equal to the bulk concentration. Therefore, the biodegradation of both substrate fractions (molecules containing \textsuperscript{12}C and \textsuperscript{13}C, short \textsuperscript{12}S and \textsuperscript{13}S) follows Michaelis-Menten kinetics\textsuperscript{42} and is described by the set of equations (2) and (3):\textsuperscript{16}

\[
\frac{d[\textsuperscript{12}S]}{dt} = \frac{q_{\text{max}}[\textsuperscript{12}S]}{[\textsuperscript{12}S] + [\textsuperscript{13}S] + K_M}
\]
where $\alpha$ is the fractionation factor with $\epsilon = \alpha - 1$, $q_{\text{max}}$ is the maximum degradation rate, and $K_M$ is the half saturation constant of the Michaelis-Menten kinetics. In the presence of mass transfer limitations across the cell envelope it is necessary to distinguish between substrate concentrations outside the cell, $S$, and substrate concentrations inside the cell, $S(\text{bio})$, where the exchange rate between these two phases is determined by the mass-transfer coefficient $k_{tr}$. Including the mass transfer limiting term in equation (2) and (3) gives equations (4) and (5) and analogous equations for the heavy fraction, where the last term in equation (5) is multiplied by the fractionation factor $\alpha$:

\[
\frac{d[^{13}S]}{dt} = \frac{\alpha q_{\text{max}}[^{13}S]}{[^{12}S] + [^{13}S] + K_M} \tag{3}
\]

These equations were solved and fitted to the experimental results to obtain the unknown parameters $k_{tr}$ and the maximum degradation rate $q_{\text{max}}$ using a modified version of ReKinSim. The enzymatic fractionation factor $\alpha = \epsilon + 1$ was determined by the fit of the Rayleigh equation (Eq. (1), Table 1) and the value for $K_M = 19 \, \mu\text{mol/L}$ was taken from the literature. Equation (6) calculates an estimate of the atrazine diffusion coefficient in lipids, $D_{\text{lip}}$:

\[
D_{\text{lip}} = \frac{k_{tr} \times \delta \times V}{A \times K_{\text{tip-w}}} \tag{6}
\]

where $\delta = 10 \, \text{nm}$ is the thickness of two lipid double membranes (5 nm each) to mimic the Gram-negative cell wall and $V = 0.5 \, \text{L}$ the volume of the cell suspension. $A$ is the bacterial total surface area calculated from an estimate of $4 \times 10^7 \, \text{cells} \cdot \text{mL}^{-1}$ (derived from the OD$_{600} =$...
0.05 with $8 \cdot 10^8$ cells·mL$^{-1}$·OD$_{600}$) and an average bacterial surface of 4 µm$^2$. $K_{lipw} = 741$ is the lipid-water distribution coefficient of atrazine.$^{46}$

**Results and discussion**

**Atrazine degradation with Gram-negative *Polaromonas sp.* Nea-C induced smaller isotope fractionation than observed with TrzN.**

Resting cells of Gram-negative *Polaromonas sp.* Nea-C with a OD$_{600} = 0.05$ degraded 30 mg/L atrazine within 24 h (see **SI Figure S1**). TrzN-catalyzed atrazine hydrolysis to hydroxyatrazine led to considerable isotope fractionation both for carbon (enrichment of $^{13}$C relative to $^{12}$C corresponding to a normal isotope effect) and for nitrogen (depletion of $^{15}$N relative to $^{14}$N representing an inverse isotope effect) (**Figure 1**). This inverse nitrogen isotope effect is characteristic of proton-assisted hydrolysis in the transition state of TrzN.$^{38}$ The enrichment factors for carbon $\varepsilon$(C) = $-3.5\% \pm 0.1\%$ and nitrogen $\varepsilon$(N) = $1.9\% \pm 0.1\%$ ($ \pm 95 \%$ confidence intervals) were determined by the Rayleigh equation, as shown in **Figure 2A, B**. These enrichment factors are significantly smaller than those described for atrazine hydrolysis catalyzed by TrzN, i.e. those obtained during biodegradation with Gram-positive *A. aurescens* TC1$^{32}$ (**Figure 2A, B**). We screened for genes analogous to the trzN gene sequence from *A. aurescens* TC1$^{47}$ in the NCBI database by Blast search$^{48}$ and found more than 20 sequences coding for TrzN with more than 99 % similarity (**SI Table S1**). Also the isotope fractionation of the abiotic model reaction – acid-catalyzed hydrolysis in water – is stronger than observed in our experiment with *Polaromonas sp.* Nea-C.$^{32}$ Taken together, this evidence is consistent with the hypothesis that a different rate determining step – mass transfer across the cell envelope – partially masked the isotope fractionation of the enzyme in *Polaromonas sp.* Nea-C. Remarkably, the enrichment factors for the Gram-negative *Chelatobacter heintzii* ($\varepsilon$(C) = $-3.7 \pm 0.2 \%$ and $\varepsilon$(N) = $2.3 \pm 0.4 \%$) are statistically indistinguishable from *Polaromonas sp.* Nea-C. Both are Gram-negative bacteria with the
difference that *Chelatobacter heintzii* degrades atrazine with a different enzyme - AtzA - but still via the same acidic hydrolysis (Table 1). This raises the question whether these different enzymes from different species (AtzA, TrzN from *Polaromonas sp.* Nea-C, and TrzN from *A. aurescens* TC1) have different transition states and thus different enrichment factors, or whether the difference in isotope fractionation is attributable to physiological differences in the cell envelope that are characteristic of Gram-positive (*A. aurescens* TC1) versus Gram-negative (*Chelatobacter heintzii* and *Polaromonas sp.* Nea-C) bacterial strains.

**Strong enzymatic isotope fractionation was masked by mass transfer limitations.**

Indeed, although the sequences of *trzN* genes from *Polaromonas sp.* Nea-C versus *A. aurescens* TC1 are highly similar (see above), it cannot be strictly excluded that subtle differences in the protein structure of even TrzN could be responsible for the differences in isotope fractionation. For example, in a recent study by Schürner et al., we observed that single point mutations in the *trzN* gene can lead to subtle changes in isotope fractionation. We therefore prepared cell free extracts of *Polaromonas sp.* Nea-C to degrade atrazine to hydroxyatrazine and followed the degradation with CSIA. As the hydrolysis of the atrazine C-Cl bond does not depend on any cofactors or energy in the form of ATP, the cell free extracts were highly active, atrazine turnover was fast (*SI Figure S2*), and was accompanied by strong isotope fractionation, as shown in **Figure 1**. The isotope fractionation in cell free extracts was considerably larger than in whole cells and gave enrichment factors $\varepsilon(C) = -5.3 \pm 0.7 \text{‰}$ and $\varepsilon(N) = 3.2 \pm 0.5 \text{‰}$ that again were indistinguishable from those of *A. aurescens* TC1 (**Figure 2 A, B**). This isotope fractionation was also similar to that of recombinant TrzN from *A. aurescens* TC1 (instead of cell free extracts), and of abiotic acidic hydrolysis (Table 1). Further, the slope of the dual element isotope plot $\lambda = \varepsilon(N)/\varepsilon(C)$ was the same for the degradation with *Polaromonas sp.* Nea-C, *A. aurescens* TC1, and the cell free extract of *Polaromonas sp.* Nea-C ($\lambda = -0.55 \pm 0.04, -0.60 \pm 0.02, \text{and } -0.61 \pm 0.14$ respectively)
(Figure 2 C) and was similar to those with recombinant TrzN and abiotic acidic hydrolysis (Table 1).\textsuperscript{32, 38} This similarity in intrinsic isotope fractionation strongly suggests that the same enzymatic reaction and same transition state prevailed but that this isotope fractionation was partially masked by a non-isotope fractionating step. This masking occurred only in whole cells with intact cell envelope, but not in cell free extracts of \textit{Polaromonas sp. Nea-C}. Such masking effects have previously been invoked to be attributable to (i) artificial high cell densities\textsuperscript{49}, (ii) diffusion through water, or (iii) retention in extracellular polymeric substance (EPS). These alternative explanations can be ruled out, however, since (i) our cell densities were small in comparison with Kampara \textit{et al.}'s study\textsuperscript{49}, (ii) the atrazine diffusion in water is fast compared to diffusion in lipid membranes,\textsuperscript{31} and (iii) \textit{Polaromonas sp. NeaC} does not form EPS. Consequently, we conclude that it was mass transfer across the cell envelope that was the partially rate-limiting step in biodegradation of atrazine by the Gram-negative \textit{Polaromonas sp. Nea-C}, but not by the Gram-positive \textit{A. aurescens} TC1.

**Passive processes dominate atrazine uptake into the cell.**

The phenomenon that organic pollutant uptake can mask isotope fractionation has already been described by Qiu \textit{et al.} where active transport along the proton motive force was the rate-determining step for phenoxy acid degradation at high concentrations.\textsuperscript{27} However, no specific transporters for atrazine are known and non-polar molecules like atrazine with a relatively high log P value of 2.6 can even permeate the phospholipid bilayer directly.\textsuperscript{50, 51} A phosphotransferase uptake system\textsuperscript{52} can be ruled out, as atrazine does not undergo phosphorylation. Other plausible uptake pathways are active transport across the lipid bilayer driven by ATP hydrolysis or by an electrochemical gradient.\textsuperscript{53-55} To explore these hypotheses, atrazine degradation rates with \textit{Polaromonas sp. Nea-C} and \textit{A. aurescens} TC1, were compared for both strains with and without addition of KCN. Cyanide is known to inhibit cytochrome c so that the proton gradient collapses and ATP production ceases. As shown in
Figure 3, the initial atrazine degradation rates in Polaromonas sp. Nea-C and A. aurecens TC1 were not influenced by 0.25 mM KCN. We conclude that atrazine degradation does not depend on active transport by ATP or the proton motive force. Thus, passive processes driven by the atrazine gradient led to atrazine uptake, e.g. through facilitated transport with porins or permeation of the membrane itself.\textsuperscript{51, 56} Note that we did not study isotope fractionation here, because (i) this concentration range was also covered in the previous degradation (SI Figure S1) and (ii) the isotope fractionation was not concentration dependent in this concentration range (Figure 2A, B; Table 1) so that the same isotopic enrichment factor is expected.

Implications for the application of CSIA in field studies.

When mass transfer masks the enzymatic reaction, this does not only limit biodegradation in the environment, but also has implications for the in situ assessment of biodegradation based on CSIA: pollutant turnover via the Rayleigh equation can best be estimated if isotope enrichment factors associated with a certain degradation pathway are constant and show little variation. However, as demonstrated in this study and by Renpenning et al.,\textsuperscript{28} the isotope fractionation does not only depend on the reaction mechanism, but also on masking of the enzymatic reaction by mass transfer across the Gram-negative cell envelope. This leads to subtle differences in enrichment factors even for the same enzymatic reaction and introduces a small additional uncertainty in biodegradation assessments by CSIA in the field.\textsuperscript{57} As a consequence, identification of the primary degradation pathway and the primary degrading strain would help to relate isotope fractionation in the field to isotope fractionation in the lab. A different strategy would be to use the enrichment factor determined under optimal conditions where mass transfer limitations are absent as a conservative estimate of biodegradation. This may underestimate biodegradation when the mass transfer becomes more and more rate limiting at low concentrations.\textsuperscript{31} In contrast, the possibility to distinguish different processes and reaction pathways with dual element isotope plots remains valid, as
long as the mass transfer across the cell envelope does not mask the enzymatic isotope fractionation completely.

Therefore, we theoretically analyzed how the isotope fractionation during atrazine degradation by *Polaromonas sp.* Nea-C is affected by decreasing concentrations. This concentration-dependent observable isotope enrichment factor $\epsilon^*$ can be modeled with a mathematical framework proposed by Thullner *et al.* for the case that mass transfer masks the intrinsic enzymatic fractionation factor $\epsilon$. This framework correlates the specific affinity of the enzyme $a = q_{\text{max}} K_M^{-1}$ with the mass transfer coefficient across the cell envelope $k_{\text{tr}}$. When the influx (determined by $k_{\text{tr}}$) is slower than the enzymatic turnover (determined by $a$) the fractionation factor $\epsilon$ will be masked which leads to a smaller observable enrichment factor $\epsilon^*$ ($\epsilon^* < \epsilon$). We used numerical modeling (see experimental section above) to fit the time-dependent enrichment in $^{13}$C associated with the atrazine concentration decrease (equations (4) and (5); SI Figure S3). Thus, we were able to estimate the parameters for Thullner *et al.*’s model: $q_{\text{max}} = 2.7 \text{ nmol L}^{-1}\text{s}^{-1}$ which gives $a = 0.14 \text{ s}^{-1}$ and the mass transfer coefficient across the cell envelope $k_{\text{tr}} = 1.6 \times 10^{-4} \text{ s}^{-1}$. We validated our modeling approach for $k_{\text{tr}}$ by calculating the lipid diffusion coefficient $D_{\text{lip}}$ according to equation (6) where the cell shape and physiology is taken into account, to compare with literature values. Indeed, the calculated atrazine lipid diffusion coefficient $D_{\text{lip}} = 1.3 \times 10^{-17} \text{m}^2\text{s}^{-1}$ was, as expected, smaller, but in the same range as $D_{\text{lip}}$ reported for atrazine in a single lipid bilayer of the Gram-positive *A. aurescens* TC1. This demonstrates, that our modeling approach yields realistic values for $k_{\text{tr}}$ allowing us to use $k_{\text{tr}}$ to predict a decreasing observed fractionation factor $\epsilon^*$ with decreasing atrazine concentrations according to Thullner *et al.*. Consistent with our experimental results, at a concentration of 4 mg/L the enzymatic fractionation factor of $\epsilon = -5.3 \%$ is already reduced to $\epsilon^* = -3.5 \%$ and it is predicted to be further reduced to below -3 % already at an atrazine concentration of 1 mg/L.
Pollutant mass transfer of non-polar pollutants may be rate limiting for biodegradation in Gram-negative bacteria.

Our results strongly suggest that the specific physiology of the Gram-negative Polaromonas sp. Nea-C with its additional restrictive outer membrane limited the influx of atrazine. In contrast, isotope fractionation in the Gram-positive A. aurescens TC1 was fully observable, demonstrating the absence of mass transfer limitation. This shows that the permeation of the cell envelope is partially rate-determining for atrazine degradation by Polaromonas sp. Nea-C already at high concentrations. Furthermore, Renpenning et al. show that the mass transfer across the cell envelope of Gram-negative bacteria affects biodegradation of chlorinated ethenes.\textsuperscript{28} Taken together, the difference between Gram-positive and Gram-negative physiology might also affect the nature of non-polar pollutant biodegradation in the environment: while a restrictive outer membrane protects Gram-negative bacteria from xenobiotics\textsuperscript{58} and from the toxicity of compounds with high log P values\textsuperscript{59, 60} it might also lower the supply of non-polar pollutants as nutrients when transporters are absent.
Figure 1. Isotope fractionation in *Polaromonas sp*. NeaC depends on the integrity of the cell envelope. The biodegradation of atrazine by the Gram-negative *Polaromonas sp*. Nea-C (black full squares) leads to considerably less isotope fractionation than the atrazine degradation of cell free extracts of *Polaromonas sp*. Nea-C (red empty squares) both for normal carbon (A) and inverse nitrogen (B) isotope fractionation. Error bars represent typical standard deviations of carbon (± 0.3 ‰) and nitrogen (± 0.5 ‰) isotope analysis.
Figure 2. Rate limiting mass transfer across the Gram-negative cell envelope was revealed by isotope fractionation. Normal carbon isotope fractionation factors ($\varepsilon(C)$) (A) and inverse nitrogen isotope fractionation factors ($\varepsilon(N)$) (B) were determined by the Rayleigh equation. Enrichment factors in cell free extracts of *Polaromonas sp.* Nea-C (red empty squares) were identical to those with whole cells of Gram-positive *A. aurescens TC1* (blue full circles) and purified TrzN$^{38}$ (green empty circles) indicating that an identical enzyme
reaction was at work. In contrast, smaller isotope fractionation was observed in degradation with intact cells of Gram-negative *Polaromonas sp.* Nea-C (black full squares). (C) The slope λ in the dual element isotope plot was similar for all degradation experiments, indicating that a common reaction mechanism (acidic hydrolysis) and similar transition state architecture is present in TrzN of both bacteria. Taken together, this indicates that the isotope effect of the enzyme reaction was partially masked by mass transfer across the cell envelope in *Polaromonas sp.* Nea-C. This non-fractionating step affects carbon and nitrogen fractionation in the same way so that the dual element isotope slope λ stays constant even though the enrichment factors are smaller. Error bars represent typical standard deviations of carbon (± 0.3 ‰) and nitrogen (± 0.5 ‰) isotope analysis (except for data from degradation with *A. aurescens* TC1 where total uncertainties are given).
Figure 3. Mass transfer of atrazine into the cytosol is not mediated by a mode of active transport that depends on energy or the proton gradient. The degradation rates for both species ((A) *A. aurescens* TC-1 and (B) *Polaromonas sp.* Nea-C) were the same for control cells (black squares) and cells treated with 0.25 mM KCN (red circles). A pseudo first order reaction kinetics was assumed, as the concentration range was well below the Michaelis-Menten constant $K_M$ of TrzN. Cyanide was added to inhibit cytochrome c to prevent formation of a proton gradient so that energy production ceases. The hydrolytic enzyme TrzN does not depend on ATP or other cofactors and is not inhibited. The degradation rates were reduced in *A. aurescens* TC-1 150 minutes after KCN addition, indicating endogenous decay of TrzN. The fits of the first order rate constant in (A) and (B) are statistically not different at the 0.05 significance level.
Table 1. Overview of isotope fractionation during atrazine degradation via acidic hydrolysis in different experimental setups. Uncertainties represent 95% confidence intervals.

| experimental system          | enzyme | Gram stain | ε(C) (‰) | ε(N) (‰) | λ = ε(N)/ε(C) | concentration (mg/L) | Ref.   |
|-----------------------------|--------|------------|----------|----------|---------------|----------------------|--------|
| whole cells Polaromonas sp. Nea-C | TrzN   | negative   | -3.5 ± 0.1 | 1.9 ± 0.1 | -0.55 ± 0.04 | 30 - 1.4             | this study |
| cell free extract Polaromonas sp. Nea-C | TrzN   | negative   | -5.3 ± 0.7 | 3.2 ± 0.5 | -0.60 ± 0.14 | 30 - 2.8             | this study |
| whole cells A. aurescens TC1 | TrzN   | positive   | -5.4 ± 0.6 | 3.3 ± 0.4 | -0.61 ± 0.02 | 18 - 1.3             | 32     |
| purified A. aurescens TC1 TrzN | TrzN   | positive   | -5.0 ± 0.2 | 2.5 ± 0.1 | -0.54 ± 0.02 | 24 - 3               | 38     |
| Chelatobacter heintzii       | AtzA   | negative   | -3.7 ± 0.2 | 2.3 ± 0.4 | -0.65 ± 0.08 | 15 - 1.8             | 32     |
| abiotic pH 3 60°C            | ---    | ---        | -4.8 ± 0.4 | 2.5 ± 0.2 | -0.52 ± 0.04 | 24 - 3               | 32     |
ASSOCIATED CONTENT

Supporting information

A more detailed experimental section, graphs of the degradation experiments, and a table of TrzN sequence similarities (PDF). This information is available free of charge via the Internet at http://pubs.acs.org.

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