The invasive crayfish *Faxonius immunis* causes the collapse of macroinvertebrate communities in Central European ponds

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Abstract The invasive crayfish *Faxonius immunis* is regarded as a threat to amphibians and macroinvertebrates in the Upper Rhine Valley, Germany, eradicating macrophytes and establishing high-density populations in stagnant waters. This study investigates the macroinvertebrate community structure of five conservation ponds south of Karlsruhe, Germany, to identify effects caused by this invasive crayfish. Two of the ponds had a high population density of *F. immunis*, two were expected to have lower crayfish densities as they had been habitat modified with gravelled water beds as a crayfish management approach, and one pond was known to be free of crayfish but contained fish. The macroinvertebrate communities were analyzed considering their species richness and composition. The relative density of *F. immunis* within the samples was regarded as a representative indicator for crayfish population density and tested for its influence on the ordination along with habitat composition and abiotic factors using distance-based redundancy analysis. *F. immunis* was identified as a driving factor of the macroinvertebrate communities of sampled ponds. Additionally, this study indicates that gravelled ground beds as a management method for invasive burrowing crayfish species does not significantly influence the community composition but can minimize indirect effects caused by *F. immunis*.

Keywords Decapoda · Management · Neobiota · Community ecology · Conservation

Introduction

Invasive non-native species are one of the biggest threats to biodiversity (Butchart et al. 2010), and invasions in freshwater habitats are of global concern (Chandra and Gerhardt 2008). They subsequently can act as driving factors in aquatic macroinvertebrate communities in rivers (Baur and Schmidlin 2008) as well as in lakes (Gergs and Rothhaupt 2015). The River Rhine itself can act as an introduction highway for aquatic invasive species (Leuven et al. 2009), and its macroinvertebrate communities are in contact with several invasive species (ICPR 2015). However, direct effects on community compositions are rarely observed, perhaps due to the complexity of the problem, which is caused by the specific ecology of invasive species (Karatayev et al. 2009). In the Upper Rhine Valley, the invasive freshwater crayfish *Faxonius immunis* was first detected in 1993 (Gelmar et al. 2006), spreading fast along the main river and oxbows...
(Chucholl 2012). By now, *F. immunis* has spread downstream to the federal state of Hesse (Herrmann et al. 2018b) with an isolated population in the city of Düesseldorf in the federal state of North Rhine-Westphalia, Germany (Gross 2019). The spread of *F. immunis* is not limited to the main river and its oxbows, and it also disperses through interconnected watercourses and ditches (Chucholl 2012; Martens 2016). The species is able to disperse over land and populate isolated ponds next to its habitat of origin at any time of the year (Herrmann et al. 2018a). This also happens for temporary ponds created for nature conservation, where the crayfish can be a threat to animals like the lilypad whiteface *Leucorrhina caudalis* or the European tree frog *Hyla arborea*, which are protected under the European Bird’s and Habitats Directive (European Economic Community 1992). *F. immunis* is known as a burrowing species (Hobbs and Hart 1959) that is defined as a tertiary burrower (Berrill and Chenoweth 1982). It can build high population densities in ponds and oxbows (Goellner 1943; Crocker and Barr 1968; Chucholl 2012), changing the habitat drastically by reducing aquatic macrophytes to a minimum and increasing turbidity of the waterbody (Martens 2016).

As the species reaches its highest densities in small ponds, it is crucial to understand the invasive potential in these locations. Studies derived from oxbows and rivers cannot reveal the role of this invader in these other highly invaded habitats. The present study compares the macroinvertebrate communities of five conservation ponds with different stages of applied habitat management and *F. immunis* population density to investigate effects caused by this invasive crayfish species and to test whether *Faxonius immunis* is a driving factor within the macroinvertebrate communities.

**Materials and methods**

**Sampling**

The sampled ponds (Table 1) were chosen based on crayfish density, comparability, accessibility and the level of gravelling the bottom sediment as management approach. They are located near the city of Rheinstetten and the community of Durmersheim, south of Karlsruhe, Germany. This area is part of the Upper Rhine Valley, which is known as a hot spot for biodiversity in Europe (Ackermann and Sachtleben 2012).

Pond A was created in 2014 as part of the EU-Life-+ Project “Rheinauen bei Rastatt” in the area Heilwald at the community of Durmersheim. It was monitored for dragonflies and for molluscs in the context of technical reports on the success of the project in 2015 (Schie 2015; Groh and Richling 2015). In the same year, pond A was described as a well-developing habitat with diverse aquatic vegetation and a good suitability as dragonfly habitat. In 2017, the pond first appeared turbid and seemed to have lost almost all of its aquatic macrophytes. It was chosen because there was a remarkably high population density of *F. immunis* known. In 2017, a population density of 10.47 individuals / m² was estimated using a mark-recapture method (own unpublished data).

The ponds B, C, D and G (Table 1) were created for threatened amphibians and used as ecological compensation for the construction of communal infrastructure and are part of an area with more than 10 ponds in the city of Rheinstetten, south-western Germany. The last changes made to the ponds had been maintenance to pond B in 2014 and the gravelling of pond C and D in the years 2017 and 2019. The conservation area in Rheinstetten is a relevant habitat of rare amphibian species like the European tree frog *Hyla arborea*, the northern crested newt *Triturus cristatus* and the agile frog *Rana dalmatina* (Beck und Partner 2014), which are listed in the EU directive 92/43EEC (European Economic Community 1992). The ponds are also habitat for different rare invertebrate species such as great silver water beetle *Hydrophilus piceus* and the lilypad whiteface *Leucorrhina caudalis* (own unpublished data).

*F. immunis* is widespread in the area. In these summer-warm ponds with mud or clay as main substrate, *F. immunis* builds high population densities. Only single ponds could be managed to have low crayfish abundances (C, D), achieved by a combination of habitat modification by gravelling the bottom sediment and reducing the population by removing crayfish over multiple years. In the context of gravelling the sediment, a part of the water surface was lost; therefore, the ponds dried out to a bigger part. Ponds B and C have had a known population of the invasive common sunfish *Lepomis gibbosus*. One
Table 1 Sampled ponds in the city of Rheinstetten and the community of Durmersheim with coordinates in decimal grades (for World Geodetic System 1984 [WGS 84]), approximate maximal depth, known fish population, expected densities of *Faxonius immunis* and their level of habitat modification

| Coordinates                  | Area [m²] | Approx. max. depth [m] | Fish                                      | *F. immunis* density | Habitat modification |
|------------------------------|-----------|------------------------|-------------------------------------------|----------------------|----------------------|
| A 48.949187, 8.281924        | 481.9     | 1.7                    | None                                      | High                 | None                 |
| B 48.973105, 8.294864        | 2918.2    | 1.6                    | *Lepomis gibbosus*                        | High                 | None                 |
| C 48.971479, 8.293772        | 2893.4    | 1.5                    | *Lepomis gibbosus*                        | Low                  | Medium               |
| D 48.971493, 8.295961        | 2115.6    | 1.0                    | None                                      | Low                  | High                 |
| G 48.970317, 8.298021        | 1441.9    | 1.8                    | *Lepomis gibbosus, Rutilus rutilus, Scardinius erythrophthalmus* | Null                 | None                 |

Single pond is free of crayfish (G) with known populations of the common rudd *Scardinius erythrophthalmus* and the common roach *Rutilus*.

Macroinvertebrates were sampled monthly in March, April and May 2019. Samples in each pond were taken using a telescopic gabion dip net (*WINLAB®, Ø 200 mm, 1.5 mm mesh size*). At the ponds, representative microhabitats were identified and partitioned to the sum of 5 microhabitats per sampling period per pond prior to sampling. Abiotic factors (temperature, pH) were recorded at the start of the sampling using a digital thermometer (GTH 1160) and pH meter (Voltcraft PH 100 ATC). The collection of macroinvertebrates consisted of ten standardized 1-m swipes per microhabitat. The retrieved material was collected in buckets, sieved over a framed 500 µm mesh and sorted alive before processing to 1-L polypropylene (PP) flasks. Amphibian larvae, bigger plant parts and stones received during the collection were put back into the water before further sorting. To prevent the collected wet material from diluting the medium too much, 96% ethanol was used for conservation in the field.

Field samples were transported to the laboratory and partitioned to bigger (e.g. juvenile crayfish, water beetles, dragonfly larvae) and smaller (e.g. chironomids, small damselfly larvae) specimens samples, and 96% ethanol was renewed prior to identification. A list of the literature used for identification is provided in supplementary material. Macroinvertebrates were counted and identified using a Leica EZ4 stereo microscope and the latest literature. Where possible, specimens were identified to species level and processed to fresh 70% ethanol. In Diptera, identification to species level was sparse and the genus- or (sub-, super-) family level was taken. The same was true for undescribed larval stages in Odonata, Coleoptera and Heteroptera.

Data analysis

Abiotic parameters and habitat compositions were carried to an explanatory factor matrix, together with corresponding levels of habitat modification. A complete list of abiotic and habitat parameters including the data type is provided in the supplements. To investigate the effect of management through habitat modification, the level of management was quantified using a 3-level scale (0; 0.5; 1). The 0.5 level describes pond D, where larger areas of the shore are covered with *Phragmites australis* and could therefore not have been gravelled completely (Table 1).

The factor identifying the relative density of *F. immunis* was created using the absolute count of *F. immunis* specimen per sample and dividing it by the total count of macroinvertebrate individuals within the same sample. The species was not taken as a response variable and therefore was removed from the macroinvertebrate dataset. The environmental factors were scaled relative to each other to values between 0 and 1 to ensure comparability using the decostand command within the package vegan 2.5–6 (Oksanen et al. 2019).
for the program R (R Core Team 2013). Each pond was tested for its absolute species richness and Shannon index, and evenness was checked for a first comparison of sites. The diversity indices taken from the Shannon index output were later analyzed towards their correlation with the density factor of F. immunis. The regression was tested using a general linear model (Gaussian family).

A dissimilarity matrix was created from the abundance data matrix. Because the macroinvertebrate dataset incorporated many rare species and the matrix therefore contained about 82% zero entries, Bray–Curtis dissimilarities were taken. To minimize the effect of outliers on the community dissimilarities of samples and to set focus on the composition of communities, the macroinvertebrate abundance data were fourth root transformed prior to analysis (Anderson et al. 2011). A hierarchical cluster analysis (McQuitty 1966) was performed prior to further analysis to test whether the sample dissimilarities matched the expected patterns.

Factors for model creation were selected using the bioenv command from vegan 2.5–6, which finds the best subset of environmental variables, so that the Euclidean distances of scaled environmental variables have the maximum correlation with macroinvertebrate community dissimilarities (Clarke and Ainsworth 1993). A distance-based redundancy analysis (db-RDA) (Legendre and Anderson 1999) was chosen to test the influence of selected factors on the macroinvertebrate communities. The significance of chosen factors was tested subsequently following the suggestions of McArdle and Anderson (2001). The received cumulative $R^2$ of the significant model, which can be displayed to high, was then corrected according to Legendre et al. (2011). Using variation partitioning (Borcard et al. 1992) by the varpart command from the vegan (3.5–6) package, the adjusted $R^2$ per explanatory factor was also calculated separately (Gherardi 2007).

## Results

### Biodiversity

In the present study, a total of 98 taxa were found within the 15 samples. Species richness was highest for pond G, where *Faxonius immunis* was not present, with 464 individuals from 52 taxa. The minimal count derived from the sample of pond A taken in May 2019 with six taxa, while the proportion of *F. immunis* within the sample was 87.34% of the individuals. Water temperatures during the study ranged from 13.4 ($\pm$ 1.1) °C in March to 21.4 ($\pm$ 2.0) °C in May and did not differ significantly between the sampled ponds. Emerging taxa abundances decreased gradually with rising water temperatures. The Shannon diversity index taken per pond showed a negative relation with *F. immunis* density (Fig. 1). The applied general linear model revealed significance of the regression between the abundance-based factor describing *F. immunis* density and decreasing species richness ($R^2 = -0.82$, $p < 0.001$).

The species richness declined from March to May, while the relative abundance of *F. immunis* increased for ponds containing the species. Pond C marks the exception with a known small population of *F. immunis*, which was not represented within the samples taken for pond C (Table 2). Ponds A and B were characterized by exceptionally low total abundances of macroinvertebrates. They contained 85% less Mollusca, 98% less Ephemeroptera larvae and 98% less Odonata larvae compared to the crayfish-free pond G (Fig. 2). For pond A, species richness declined from 22 to 5 detected species over the sampling period. The abundances of aquatic Heteroptera were highest for pond C with 148 individuals.

The complete abundance data table containing all taxa and individual counts per sample is provided in supplementary material.

### Dissimilarity

Hierarchical cluster analysis clearly separated samples of ponds taken in March and April from those sampled in May 2019 at the 90% dissimilarity level. The crayfish-free pond G clusters isolated and shows highest similarity with itself over the whole sampling period (Fig. 3). The ponds B–G show similarities within the samples from March and April.

### Community analysis

The best subset of environmental variables, as selected by the bioenv function, included water temperature (T), proportion of *Juncus* spp. and mud as microhabitat compartments and the abundance-based factor for
**Faxonius immunis** with a correlation of 69.3%. The applied db-RDA clearly showed gradient-driven ordination towards the density of *Faxonius immunis* (Fig. 4). The explained variation from the db-RDA output was 40.9% after correction of $R^2$. As the first two axes together account for 69.71% of the ordination, a 2-dimensional plot was chosen.

**Table 2** Total counts for taxa and number of individuals (without *F. immunis*) and total counts as well as relative proportion of *F. immunis* per sample in five ponds north of Karlsruhe, Germany

| Sample | Taxa | Individuals | Total | Relative |
|--------|------|-------------|-------|----------|
| **A**  | March| 22          | 68    | 1        | 0.015    |
|        | April| 21          | 45    | 27       | 0.375    |
|        | May  | 5           | 13    | 90       | 0.874    |
| **B**  | March| 38          | 184   | –        | –        |
|        | April| 28          | 66    | 20       | 0.230    |
|        | May  | 23          | 38    | 43       | 0.530    |
| **C**  | March| 14          | 171   | –        | –        |
|        | April| 26          | 274   | 2        | 0.007    |
|        | May  | 16          | 49    | 12       | 0.197    |
| **D**  | March| 14          | 171   | –        | –        |
|        | April| 14          | 317   | –        | –        |
|        | May  | 9           | 105   | –        | –        |
| **G**  | March| 27          | 93    | –        | –        |
|        | April| 23          | 130   | –        | –        |
|        | May  | 11          | 92    | –        | –        |

**Fig. 1** Shannon diversity indices during the sampling period for the sampled ponds with high population densities of *Faxonius immunis* (dark grey), managed ponds with low densities (light grey) and the crayfish-free pond (white). Whiskers marking the total range.

**Fig. 2** Number of individuals caught within the taxonomic groups of Mollusca, Ephemeroptera and Odonata larvae for high-density ponds, managed ponds with lower crayfish densities and the crayfish free pond. Error bars indicating the standard error within the data row. The row for pond G has no error bar and is only given for optical comparability.

**Fig. 3** Hierarchical cluster analysis of similarities for samples taken from the studied ponds A, B, C, D and G, in March, April and May (1, 2 and 3), respectively. Marked abbreviations highlight high ponds with high densities of *F. immunis* (dark grey), managed ponds with lower crayfish densities (light grey) and the crayfish free pond (thin line).
The variance inflation factors of the chosen subset did not show signs of multicollinearity and ranged from 1.12 to 1.79 (Table 3). The variation partitioning algorithm showed little overlap between factors. The proportion of total variance explained by each single factor was highest for *Faxonius immunis* (13.27%), followed by water temperature (12.2%) and lower for *Juncus* spp. (8.92%) and mud (6.38%) as microhabitat proportions of the ponds.

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**Discussion**

The present study clearly shows strong impacts of *Faxonius immunis* on the studied macroinvertebrate communities. The decrease in macroinvertebrate abundances, Shannon biodiversity indices and α-diversity along a crayfish-density gradient and the resulting increase in its dominance is clearly visible. Changes in macroinvertebrate community structure as well as the differences visualized in the db-RDA ordination are oriented along the density gradient as well. Previous findings reporting effects of crayfish invasions on macroinvertebrate communities in North

**Table 3** Subset of factors with their variance inflation factors (vif), biased and adjusted $R^2$ as well as p-values and degrees of freedom (F) Taken from the margin test output

| Factor          | vif   | F     | $R^2$  | adjusted $R^2$ | p value   |
|-----------------|-------|-------|--------|----------------|-----------|
| *Faxonius immunis* | 1.794 | 3.0186| 0.19425| 0.13227        | 0.001***  |
| Water temperature| 1.161 | 3.1621| 0.18465| 0.12193        | 0.001***  |
| *Juncus* spp.    | 1.125 | 3.0896| 0.15426| 0.08920        | 0.001***  |
| Mud             | 1.763 | 1.7123| 0.13070| 0.06383        | 0.090     |

*Represents significance at the alpha < 0.05 level
America (Albertson and Daniels 2016) as well as studies on effects on the endemic aquatic flora and fauna by other invasive crayfish such as *Procambarus leniusculus* in Sweden and England (Nyström et al. 1996; Mathers et al. 2016) or *Faxonius virilis* in the Netherlands (e.g., Roessnik et al. 2017) describe similar effects. Such results were summarized and seem to overlap in most invasive, non-native crayfish species (Twardochleb et al. 2013).

The results of the present study can best be compared to documented effects of *Procambarus clarkii* in Spain, Portugal or Italy (Gherardi and Acquistapace 2007; Rodríguez et al. 2005; Correira and Anastácio 2008; Barbaresi and Gherardi 2000) and underline suggested invader-specific ecological impacts of *F. immunis* as pronounced by Chucholl and Chucholl (2021). The relations between *F. immunis* density and α-diversity outline the relevance of ongoing crayfish management in areas where rare macroinvertebrate species occur. With its strongly r-selected life history (Tack 1941; Chucholl 2012) and its ability to migrate over land (Herrmann et al. 2018a), *F. immunis* will be a growing threat for macroinvertebrate communities and ecosystem services along waterbodies invaded by this crayfish species.

The macroinvertebrate communities from ponds with high *F. immunis* densities in the present study were characterized by very low abundances in macrophyte-associated taxa, especially Odonata larvae and Mollusca. Even Ephemeroptera larvae, such as *Cloeon dipterum*, a species which was expected to be unaffected by the crayfish, had very low abundances in samples from high-density ponds. The effect on Odonata and Mollusca was most radical in pond A, which was monitored for dragonflies and molluscs in 2015, one year after its creation. It was documented as a very suitable habitat for dragonflies with eleven species present within the samples from pond A. In contrast to the seven recorded aquatic mollusc species in 2015 (Groh and Richling 2015), none was found again in the present study. It is known that habitat structure and complexity has mediating effects on predatory taxa (Warfe and Barmuta 2004), such as odonata larvae. Although there are known negative effects on aquatic snails caused by crayfish predation (Nyström and Pérez 1998), no such population collapse was documented. However, Holdich (1999) suggests that total collapses of taxa can be caused by established high populations of invasive crayfish. Enclosure experiments also showed eradication of snails by an invasive non-native crayfish species (Chucholl 2013).

Incomplete sampling or effects on sample composition caused by the sampling method should not have played a major role in this study. García-Criado and Trigal (2005) compared sampling techniques for pond macroinvertebrate communities and found that 1-m swipes, as used in this study, give the most complete view on the macroinvertebrate communities in sampled ponds. Merolimnic species were abundant in all samples taken in March and April. Therefore we explain the increasing differences from March to May between the macroinvertebrate community samples, namely a drastic reduction in α-diversity and total abundances, with the increasing activity of *F. immunis* adults with rising water temperatures in spring (Crawshaw 1974) and the presence of newly hatched juvenile crayfish in April and May (Tack 1941; Chucholl 2012).

As crayfish can act as ecosystem engineers (Statzner et al. 2000; 2002), the effects of their invasion are not caused by predation alone but also by indirect methods (Momot 1995; Nyström et al. 1996). *F. immunis* can cause indirect effects linked to its abundance through changes in habitat structure, as the species is known to reduce macrophytes to a minimum (Tack 1941; Letson and Makarewicz 1994) and is described as a tertiary burrowing species (Hobbs and Hart 1959; Crocker and Barr 1968; Berrill and Chenoweth 1982) that increases the turbidity of the waterbody (Martens 2016). In the present study, the effects on the habitat caused by high densities of *F. immunis* are clearly visible in pond A, where the only cover left for macrophyte-associated taxa like Odonata or Coleoptera larvae is stocks of *Juncus* spp. and *Phragmites australis* at the edge of the littoral zone with the major habitat compartment consisting of muddy ground. As aquatic macrophytes and Characeae are reduced by *F. immunis*, mud becomes the dominant microhabitat structure in ponds with high crayfish densities. The increase in muddy character of a pond and the reduction of aquatic macrophytes is therefore one of the main indirect effects caused by this invasive crayfish. Roessnik et al. (2017) described negative effects on aquatic macrophyte abundances...
linked to the density of *Faxonius virilis*, an invasive species in the Netherlands. In their study, crayfish significantly reduced macrophytes starting from a crayfish density of 1.25 crayfish per m². Although mud lost its significant p-value in the present study after variation partitioning, it was left in the ordination because it is one of the main indirect effects of *F. immunis*.

Following this change in habitat composition, *F. immunis* indirectly changes the macroinvertebrate community composition in favour of mobile and pioneer species with lesser need for cover, while abundances of macrophyte-associated species and slow-moving taxa decrease. This effect was also described for the invasive crayfish *Pacifastacus leniusculus* in Sweden, where ponds with higher crayfish density inhabited mostly ground dwelling species such as Chironomid larvae or mobile taxa like aquatic Heteroptera (Nyström et al. 1996). Negative effects on the macroinvertebrate communities by predation and changes in food webs caused by *Procambarus clarkii* are frequently described and reviewed by Souty-Grosset et al. (2016). Shallow, fish free ponds in Central Europe are normally free of crayfish. The communities associated with this habitat therefore could be seen as naïve to omnivorous crayfish, as the top predators in the pond communities are dragonfly and water beetle larvae. This might intensify the effect on the community as well.

Gravelling the ground beds to manage *F. immunis* in ponds does not reduce the crayfish density itself, but the effects on habitat composition can be largely negated. While macrophytes density might still decrease, there is cover for macroinvertebrates as well as amphibian larvae between the gravel stones. Similar effects are known for rocks of different size that are deterring crayfish predation on fish larvae (Savino and Miller 1991). The increase in turbidity and muddy character of ponds is weakened in ponds with gravelled beds (Borner et al. 1997), which seems to counteract the main indirect effects caused by *F. immunis* on macroinvertebrate community structure in the present study as the species prefers muddy grounds (Caldwell and Bovbjerg 1969; Bovbjerg 1970). This management approach does not significantly alter the community composition itself, as the cluster analysis revealed seasonal clustering of samples taken in March and April for the ponds B, C, D and G, respectively. Therefore, the differences occur mainly with rising activity and the increasing number of juveniles of *F. immunis*.

*Faxonius immunis* significantly alters the macroinvertebrate community and strongly changes the habitat composition in stagnant waters by predation and otherwise reducing macrophytes. Larvae of threatened and rare macroinvertebrate species, herbivores or aquatic snails are no longer able to find the needed refuge, prey or grazing substrate between submerged macrophytes in habitats colonized by *F. immunis*. Only mobile and pioneer species seem to sustain in areas where this invasive crayfish species has built high densities. Without applied management, habitats invaded by *F. immunis* act as traps for macroinvertebrates trying to reproduce and are lost for conservation purposes.

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**Authors’ contributions** All authors contributed to the study conception and design. Final conception, field sampling and data analysis were performed by Alexander Herrmann. Identification was done by Alexander Herrmann with support from Karsten Grabow and Andreas Martens. The first draft of the manuscript was written by Alexander Herrmann, and all authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.

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**Data availability** The environmental dataset and species lists for the sampled ponds will be provided as supplementary material. All raw data and the conserved macroinvertebrate samples are available at the institute and can be viewed on reasonable request.

**Code availability** For analysis, the Software R Studio was used. The custom R code used for analysis is available from the corresponding author under the provided e-mail address.

**Declarations**

**Conflict of interest** The authors declare that they have no conflict of interest to declare.

**Ethical approval** All animals were handled according to the German laws and ethics. The local administrations and responsible officials were informed and approved the study. No specific ethics approval was needed.
Consent to participate  All authors and persons named in acknowledgements section are in consent about the manuscript submission in its actual state.

Consent for publication  All authors are in consent that the present results should be published.

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