Detection of *Ramularia collo-cygni* DMI- and SDHI-resistant field populations in Austria and the effect of fungicides on the population and genetic diversity

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**Abstract** Winter barley (*Hordeum vulgare* L.) is the third most cultivated crop after corn and wheat in Austria but one of the most challenging for disease control. The foliar pathogen *Ramularia collo-cygni* B. Sutton and J.M. Waller, causing Ramularia leaf spots (RLS), is one of the most important diseases in barley. In the recent years, control has only been achieved using fungicide mixtures including the multi-site inhibitor chlorothalonil, however this compound is totally banned in the EU. The objective of this study was to assess fungicide dose-rates and spray mixtures for RLS control. Furthermore, a field monitoring within the main barley growing areas of Austria was carried out, to analyse the current resistance situation to DMI and SDHI fungicides, which are still the backbone in RLS control. The results indicate that only the mixture with chlorothalonil achieved a good RLS control. Prothioconazole or benzovindiflupyr (alone or additively) decrease the severity of RLS but increase the local frequency of *Cyp51* and *sdhC* mutations, especially the high dose rates. Based on a low *Cyp51* mutation frequency of 16% in untreated control this frequency increased over 3.8 times following an application with 300 g ha$^{-1}$ prothioconazole. The cumulative-*sdhC* mutations were even more increased after an application with benzovindiflupyr. This study showed that *Ramularia collo-cygni* is present in 91% of barley fields presented in this field survey. Widespread use of chlorothalonil fungicide maintained a low to moderate mutation frequency (*Cyp51-I325T*, *Cyp51-I328L*, *sdhC-H146R* and *sdhC-H153R*) in Austrian barley regions with no increase between 2017 and 2019.

**Keywords** *Ramularia collo-cygni* · Fungicide resistance · Demethylation inhibitors · Succinate dehydrogenase inhibitors
Introduction

Over the last years, Ramularia leaf spot (RLS) disease has become one of the most important threats in barley (Hordeum vulgare L.). The causal agent of this late season disease is Ramularia collo-cygni B. Sutton and J.M. Waller which is an important pathogen of barley across many climates (Havis et al., 2015; McGrann & Havis, 2017) and has been detected in barley samples worldwide (Stam et al., 2019). The fungal pathogen causes necrotic leaf spots with the awns and grains also affected. The disease rapidly increases after flowering, the infection is also accompanied by premature senescence, loss of green leaf area (GLA) and substantial yield losses of around 25% (Harvey, 2002). In addition, barley is also susceptible to other foliar diseases such as powdery mildew of barley (Blumeria graminis f. sp. hordei), net blotch (Pyrenophora teres), barley scald (Rhynchosporium secalis) and leaf rust of barley (Puccinia hordei).

The main management strategies against these fungal diseases are focused on crop rotations, resistant or tolerant varieties in combination with fungicide treatments (Hess et al., 2009). In the recent years, numerous successes have been realized in breeding new varieties which are resistant to some of these diseases: e.g. mildew resistance locus 0 (MLO) genes are used in breeding new varieties for powdery mildew of barley. In Blumeria graminis f. sp. hordei, net blotch (Pyrenophora teres), barley scald (Rhynchosporium secalis) and leaf rust of barley (Puccinia hordei). The main management strategies against these fungal diseases are focused on crop rotations, resistant or tolerant varieties in combination with fungicide treatments (Hess et al., 2009).

In the European market, fungicides are subject to continuous change and will be strongly limited in the future. Chlorothalonil—a multi-site inhibitor which is a very effective chemical in control of RLS (Havis et al., 2020)—had the latest EU date of use May 20, 2020 (AGES 2020). Therefore, new fungicide strategies for RLS control will be required, especially in the light of fungicide resistance. Resistance to fungicides will be favoured especially when fungicides with a single site of action are applied (Brent & Hollomon, 2007). This is also reflected in R. collo-cygni, which showed fast adaptability to quinone-outside inhibitors (QoIs), demethylation inhibitors (DMIs) and succinate dehydrogenase inhibitors (SDHIs), the main group of fungicides used in barley crops (Rehfus et al., 2019). Fungicides based on QoIs (strobilurins, e.g. azoxystrobin) were first launched in 1996 and provided initially good control of RLS, however resistance quickly developed. A point mutation at codon 143, G143A in the cytochrome b gene in R. collo-cygni gives full resistance to QoI fungicides (Leadbeater, 2012). In another study, 93% of tested isolates were highly resistant to strobilurins in 2015 (Matsuzaki et al., 2019). This underlines the fact that a quick development of strobilurin resistance resulting in a full resistance in R. collo-cygni within 2 years had taken place (Fountaine & Fraaije, 2009).

The triazolinthione derivate prothioconazole is a systemic fungicide and belongs to the group of dem ethylation inhibitors (DMIs). The widely used curative and protective fungicide inhibits the 14α-sterol demethylation enzyme (CYP51) and leads to an inhibition of fungal growth (Parker et al., 2011; Stammmer & Semar, 2011). Compared to other single-site inhibitors such as QoIs or SDHIs, resistance development to DMIs is not so fast and substances within the group are slightly different in the development speed and selectivity of resistance (Fraaije et al., 2007). In recent years, reduced sensitivity to the DMIs have already been reported for the wheat pathogen Zymoseptoria tritici (Heick et al., 2017a, 2017b; Leroux & Walker, 2011). In the study by Rehfus et al. (2019), 12 different amino acid alterations in the coding region of the CYP51 enzyme of R. collo-cygni were found. Here, the dominant haplotype was RccC1, which contains the mutations Cyp51-I381T, Cyp51-I384L and Cyp51-Y459C. After a re-numbering of amino acids, the two mutations Cyp51-I381T and Cyp51-I384L are homologous to Cyp51-I325T and Cyp51-I328L, which
occurs always in combination (Torriani S. 2020, personal communication).

The group of succinate dehydrogenase inhibitors (SDHIs) is one of the fastest growing fungicide class, which inhibits the succinate dehydrogenase (SDH) in the mitochondrial respiratory chain. The newer generations of SDHIs, e.g. benzovindiflupyr, have a broad-spectrum of fungal activity, including *R. collo-cygni* (Avenot & Michailides, 2010; Sierotzki & Scalliet, 2013). Benzovindiflupyr is a preventive fungicide and due to the reduced stereo-centres in the structure when compared to others, e.g. isopyrazam, linked with excellent crop tolerance. An additional physiological effect is a reduced transpiration without negatively affecting photosynthesis. This results in an increase in biomass under water saving conditions (Kuznetsov et al., 2017).

Resistance to SDH fungicides have been reported after a few years of use — first in barley against *P. teres* or *R. collo-cygni* and later in several other pathogens e.g. *Z. tritici* and *Puccinia striiformis* in wheat (Cook et al., 2021; Piotrowska et al., 2017a). The SDH enzyme is composed of four nuclear-encoded subunits (sdh-A, sdh-B, sdh-C and sdh-D), whereby subunits sdh-B, sdh-C and sdh-D interact with SDHI fungicides, which can result in amino acid substitutions (Gutiérrez-Alonso et al., 2017; Sierotzki & Scalliet, 2013). Most mutations with amino acid substitutions were shown to be in the ubiquinone-binding site sdh-B, with histidine (H) most often replaced with tyrosine (Y) (Sierotzki & Scalliet, 2013). In the *R. collo-cygni* mutations *sdhC*-H146R and *sdhC*-H153R, a histidine at position 146 or 153 were substituted with arginine (R). The mutation *sdhC*-H146R (former notation *sdhC*-H142R) is homologous to mutation *sdhC*-H134R of *P. teres* and *Alternaria* species as well as to the mutation *sdhC*-H146R of *Sclerotinia sclerotiorum*. Mutation *sdhC*-H153R (former notation *sdhC*-H149R) is homologous to mutation *sdhC*-H152R and *sdhC*-H151R of *Z. tritici* and *Venturia inaequalis* (Torriani et al., 2017). Both mutations and additionally the amino acid alterations *sdhC*-H146L, *C-G91R* and *C-G171D* are associated with a significantly decrease in sensitivity to SDHIs. Furthermore, mutations *sdhC*-N87S, *B-T267I* and *B-N224T* are linked to a lower resistance factor and not as relevant as the above-mentioned mutations (FRAC 2021). Monitoring programmes in Europe also found mutations *sdhB*-264P, *B-266R/Y/L*, *B-1268V*, and *C-R152M* which have an impact on SDH sensitivity (Rehfus et al., 2019).

Due to the economic importance of *R. collo-cygni* and the lack of new active fungicidal compounds against this disease, current fungicides must be used carefully and accordance with strict guidelines to ensure their activity is maintained long as possible. To date, few sound field studies on the effect of different fungicides strategies against RLS in populations with DMI- and SDHI resistant mutations have been conducted and only limited data is available highlighting the current frequencies of *Cyp51* and *sdhC* mutations in populations of *R. collo-cygni* (FRAC 2021; FRAC 2019; Piotrowska et al., 2017a, 2017b; Rehfus et al., 2019; Torriani et al., 2017). In 2016 and 2017 first strains resistant to DMI and SDHI were found in Austria (FRAC 2021; FRAC 2019). The low to moderate DMI and SDHI resistance alleles in Austrian *R. collo-cygni* populations were used as starting point for our investigation. The purpose of this study was (I) to provide an overview about the frequency and distribution of selected *Cyp51* and *sdhC* mutations in *R. collo-cygni* populations in the main Austrian barley growing areas, (II) to assess the effect of fungicide doses (prothioconazole, benzovindiflupyr and chlorothalonil) on *Cyp51* and *sdhC* alterations and (III) to evaluate alternative fungicides to enable a replacement strategy without chlorothalonil.

**Materials and methods**

**Experimental design**

Field trials were conducted in the growing season 2017/2018 and 2018/2019, respectively. Both years were considered the warmest in the region since 1961 (ZAMG 2020). The trials were arranged as randomized complete block design with four replicates and a plot size of 15 m². All fungicides were applied using a plot sprayer in 300 L ha⁻¹ water at 2.5 bar pressure, with flat fan venturi nozzles (Albutz AVI 80–015), spaced at 25 cm on the...
sprayer boom. The distance between sprayer boom and the barley plants were kept at 25 cm.

Fungicide dose rate trial

During the growing season 2017/2018, two field trials were carried out in Wels (Upper Austria, Lat 48.20077, Long 14.04151) and St. Poelten (Lower Austria, Lat 48.20618, Long 15.59586). Both trials were located in the main Austrian barley growing areas. The soil type in Wels was a silty loam and the soil type in St. Poelten was a silty clay loam. Local two-row winter barley varieties known for their susceptibility to *R. collo-cygni* were sown in the first week of October 2017 (Wels, cv. SU Vireni; St. Poelten, cv. Sandra). Four commercially available fungicides belonging to three different modes of action and different classes of chemical were used: Proline (prothioconazole; P; Bayer Crop Science, Monheim am Rhein, Germany), Elatus Plus (benzovindiflupyr; B; Syngenta Crop Protection, Basel, Switzerland), Elatus Era (prothioconazole plus benzovindiflupyr; PB; Syngenta Crop Protection, Basel, Switzerland) and Balear (chlorothalonil; FMC Corporation, Philadelphia, USA; tank mixed with Elatus Era; PBC; (Table 1). At both sites fungicide applications were performed on May 1, 2018 at growth stage (GS) 49 (43–49) and harvest was carried out on June 20, 2018 at Wels and on June 21, 2018 at St. Poelten. From each plot, 15 random leaves of the upper two leaf-layer, were collected at GS 87, dried at room temperature for one day and frozen at -20 °C until further use.

Alternative management strategies

Three field trials were completed in the growing season 2018/2019 to evaluate alternative fungicide management strategies in Kirchdorf am Inn (Upper Austria, Lat 48.296444, Long 13.304712), Marchtrenk (Upper Austria, Lat 48.214329, Long 14.143009) and St. Poelten (Lower Austria, Lat 48.20795, Long 15.59441). For all sites soil type was silt loam. Winter barley (Kirchdorf cv. SU Vireni; Marchtrenk cv. Finola; St. Poelten cv. Anemone) was sown between September 26 and October 1, 2018 and harvest was carried out on July 3, 2019. The fungicide management strategy was developed to protect the barley crop against the main barley diseases. Thus, all plots, except the untreated control, received an early fungicide application at GS 31–32 to protect barley against *Oculimacula yallundae, B. graminis, P. teres, R. collo-cygni*.

Table 1 Fungicide treatments and total amount of active ingredients (a.i.) applied in dose-rate trial

| Treatment*,** | Product name, dose-rate [l ha\(^{-1}\)]*** | Active ingredient [g ha\(^{-1}\)] |
|--------------|---------------------------------|----------------------------------|
| **UTC**      | -                               | Prothioconazole: 87.5, Benzovindiflupyr: 87.5, Chlorothalonil: 87.5 |
| P25          | Proline, 0.15                    | 38                               |
| P50          | Proline, 0.30                    | 75                               |
| P100         | Proline, 0.60                    | 150                              |
| P200         | Proline, 1.20                    | 300                              |
| B25          | Elatus Plus, 0.19                | 19                               |
| B50          | Elatus Plus, 0.38                | 38                               |
| B100         | Elatus Plus, 0.75                | 75                               |
| B200         | Elatus Plus, 1.50                | 150                              |
| PB25         | Elatus Era, 0.25                 | 38                               |
| PB50         | Elatus Era, 0.50                 | 75                               |
| PB100        | Elatus Era, 1.00                 | 150                              |
| PB200        | Elatus Era, 2.00                 | 300                              |
| PBC25        | Elatus Era, 0.25 + Balear, 0.25  | 38                               |
| PBC50        | Elatus Era, 0.50 + Balear, 0.50  | 75                               |
| PBC100       | Elatus Era, 1.00 + Balear, 1.00  | 150                              |
| PBC200       | Elatus Era, 2.00 + Balear, 2.00  | 300                              |

*UTC* = untreated control; P = prothioconazole; B = benzovindiflupyr; PB = prothioconazole plus benzovindiflupyr; C = chlorothalonil

**Numbers stand for the application rates: 25%, 50%, 100% and 200% of the recommended rate

*** Manufacturer:

Proline = Bayer Crop Science; Elatus Plus and Elatus Era = Syngenta Agro GmbH; Balear = FMC Agro Austria GmbH
*secalis* and *P. hordei* between April 9 and April 11, 2019. The main application against RLS was performed at GS 39–49 (Kirchdorf May 6; Marchtrenk May 7; St. Poelten May 2) and GS 49–59 (Kirchdorf and Marchtrenk May 16; St. Poelten May 17) according to Table 2.

Disease scoring, green leaf area and yield assessment

Disease was assessed using EPPO protocol (Anon. 2012). Visual disease assessments for *R. collo-cygni* were carried out at four random positions per plot. The percentage diseased leaf area (disease severity, Table 2 Fungicide treatments, application timings, dose rates and the total amount of active ingredients (a.i.) applied in trials with alternative management strategies

| Treatment | GS 31–32 Dose | GS 39–49 Dose | GS 49–59 Dose |
|-----------|----------------|----------------|----------------|
| L/kg ha$^{-1}$ | a.i. (g) | L/kg ha$^{-1}$ | a.i. (g) | L/kg ha$^{-1}$ | a.i. (g) |
| 1 UTC* | | | |
| 2 Cyprodinil + Metconazole | 0.5 + 1.0 | 375/60 | Prothioconazole/Benzovindiflupyr | 1.0 | 150/75 |
| 3 Cyprodinil + Metconazole | 0.5 + 1.0 | 375/60 | Prothioconazole/Benzovindiflupyr + Chlorothalonil | 1.0 + 1.2 | 150/75/600 |
| 4 Cyprodinil + Metconazole | 0.5 + 1.0 | 375/60 | Prothioconazole/Benzovindiflupyr + Sulfur | 1.0 + 6.0 | 150/75/4800 |
| 5 Cyprodinil + Metconazole | 0.5 + 1.0 | 375/60 | Prothioconazole/Benzovindiflupyr + Folpet + Adjuvant | 1.0 + 1.5 + 0.125 | 150/75/750/80 |
| 6 Cyprodinil + Metconazole | 0.5 + 1.0 | 375/60 | Prothioconazole/Benzovindiflupyr + Folpet + Azoxystrobin | 1.0 + 1.5 + 1.5 | 150/75/750/125 |
| 7 Cyprodinil + Metconazole | 0.5 + 1.0 | 375/60 | Prothioconazole/Benzovindiflupyr | 1.0 | 150/75 | Folpet | 1.5 | 750 |

Active Ingredients
- Cyprodinil 750 g kg$^{-1}$
- Metconazole 120 g L$^{-1}$
- Prothioconazole 150 g L$^{-1}$ + Benzovindiflupyr 75 g L$^{-1}$
- Chlorothalonil 500 g L$^{-1}$
- Sulfur 800 g kg$^{-1}$
- Folpet 500 g L$^{-1}$
- Adjuvant 640 g L$^{-1}$ (= organosilicone s. fluid 385 g L$^{-1}$ + synthetic latex 255 g L$^{-1}$)
- Azoxystrobin 250 g L$^{-1}$

Product name
- Unix®
- Plexeo™
- Elatus™ Era
- Bravo® 500
- Thiovit® Jet
- Folpan® 500 SC
- Designer®
- Amistar®

Company
- Syngenta (Switzerland)
- Syngenta (Switzerland)
- Syngenta (Switzerland)
- Syngenta (Switzerland)
- Syngenta (Switzerland)
- Syngenta (Switzerland)
- Adama (Israel)
- Nufarm (Australia)
- Syngenta (Switzerland)

*UTC = untreated control
DS %) on flag leaf (F) and second leaf (F-1) was determined and the fungicide efficacy (%) was calculated using the formula (Zhang et al., 2020)

\[ \text{Efficacy} \% = \frac{A_{(utc)} - A_{(ft)}}{A_{(utc)}} \times 100 \]

where \( A_{(utc)} \) = mean lesion area in the untreated control (utc) plots and \( A_{(ft)} \) = mean lesion area in the fungicide treated (ft) plots. For the dose-rate trial the area under disease progress curve (AUDPC) (Shaner & Finney, 1977) between application and GS 87 was calculated from the RLS scores for each plot using the formula

\[ \text{AUDPC} = \sum_{i=1}^{n} \left( \frac{Y_{i+1} + Y_{i}}{2} \right) \times (X_{i+1} - X_{i}) \]

where \( Y_{i} \) = percent RLS at the ith observation, \( X_{i} \) = time in days at the ith observation, \( n \) = total number of observations, and \( Y_{0} = X_{0} = 0 \). The total number of observations were 3 in St. Poelten and 4 in Wels, respectively.

Green leaf area (GLA) was assessed similarly to disease control and the area under green leaf area curve (AUGLAC) between GS 39 and harvest was calculated using the same formula as for AUDPC whereas \( Y_{i} \) = percent of GLA at the ith observation. The total numbers of observations were 6 in St. Poelten and 7 in Wels.

Barley was harvested separately for each plot at plant maturity and grain weight were adjusted to 86% dry matter. Yield was expressed as dt ha\(^{-1}\).

Field monitoring

To get an overview about the progress of Cyp51 and sdhC alterations, a field monitoring over the main barley growing areas were carried out in Austria. Therefore, samples from fungicide free barley fields were collected. In 2017, leaves from 7 locations, in 2018 from 29 locations and in 2019 from 22 locations were analysed. 50 leaves with visible RLS per location were randomly collected, dried at room temperature for one day and frozen at -20 °C until further use. Leaves of each location were divided into two subsamples and separately analysed. The results are presented as the average frequency from the two subsamples.

Determination of Cyp51 and sdhC alterations associated with fungicide resistance

DNA from punched leaves was extracted and analysed as described by Wullschleger et al. (2015), with adjusted primers and PCR conditions (Torriani S. and Accardo S. 2020; written communication). A singleplex SYBR Green real time PCR assays using 384 well plates were carried out for detection of the quantitative mutations leading to Cyp51-I325T and Cyp51-I328L in the CYP51 enzyme and sdhC-H146R and sdhC-H153R in the sdh-C gene. The qPCR was established in Syngenta Resistance Centre in Stein, Switzerland. PCR assays were done according to the method described by Assinger et al. (2021).

Data analysis

All linear mixed models were created Proc Mixed procedure of SAS 9.4 (SAS®, SAS Institute Inc., Cary, NC). Degrees of freedom method was kr2 and covariance matrices were CS.

To analyse AUGLAC and resistance frequencies, treatments and sites were considered fixed effects, and the blocks were considered as random effect within the site. Selected data from dose-rate trials were described separately and analysed by location. In field experiments, due to unequal variances, means were compared using Hochberg’s GT2 multiple comparison test (\( p = 0.05 \)). A model for mutation presence or absence due to years and loci was created with PROC GENMOD with zero inflated negative binomial distribution. Overdispersion was confirmed and the model fit was adequate (Scaled Pearson \( \chi^2 > 0.05 \)). Map was created with spotfire 10.3 (TIBCO Software Inc., Palo Alto, CA).

To analyse AUDPC and yield, the treatments were parameterized into rate levels by fungicide and log of the standard rates were used to linearize the explanatory variables. There was little evidence of by-site interactions for AUPDC or yield; in the models presented fungicide rates and their interactions were considered fixed and blocks within sites were considered random. The fungicide active ingredients were tested individually and in their full factorial interactions by rate: prothioconazole and benzovindiflupyr were assessed individually and in interaction, and chlorothalonil was assessed only in interaction with prothioconazole and benzovindiflupyr according to the study...
design. The highest degree rate polynomial (quartic) with full factorial was fit. Visual inspection of residual plots verified there were no serious departures from model assumptions. The parameter estimates and confidence intervals are interpreted at the highest polynomial level where Type 1 sum of squares were significant at \( p < 0.05 \).

Additionally, the relationship between fungicide, AUDPC and yield was assessed through causal inference in SAS using Proc Causalmed. Normal mediator (AUDPC) and outcome (yield) distributions were assumed. Each fungicide treatment was considered individually versus the untreated control and covariates used were site, block and rate. Total effects were considered statistically significant for causal interpretation at \( p < 0.05 \).

**Results**

In both experimental years 2018 and 2019, RLS achieved between 25 and 93% infected leaf area on the upper two leaves. Lowest RLS disease severity was assessed with 25% in Marchtrenk in 2019. Highest values were measured in St. Poelten with 93% in 2018.

Effect of dose dependent fungicide treatments on disease level, green leaf area, yield and selection pressure on \( Cyp51 \)- and \( sdhC \)-mutations

All fungicide treatments significantly increased in green leaf area (except P25 in both sites and B25 in Wels) compared to the untreated control (Figs. 1A & 2A). The higher dose-rates provided greater increase in AUGLAC in St. Poelten compared to Wels. The strongest effects were obtained, when combining all three active compounds in full (PBC100) or double (PBC200) dosage. These two treatments showed significant differences in AUGLAC and additionally these plots had significant greener leaves compared to other treatments. The higher dose-rates resulted in greener leaves and were associated with a reduced RLS severity.

RLS epidemics were high at both experimental sites. In the untreated control plots RLS severity at GS 87 was 77% on flag leaf and 87% on second leaf in Wels and 91% on flag leaf and 95% on second leaf in St. Poelten, respectively an AUDPC level of 1119 for Wels and 1232 for St. Poelten. All fungicide treatments reduced AUDPC compared to the untreated control, and progressive disease reductions corresponded to dose rate (Fig. 3, Supplementary Table S1). Linear and quadratic terms for prothioconazole and benzovindiflupyr were significant, as was their first order interaction. Additionally, benzovindiflupyr had a significant interaction with prothioconazole on a quadratic basis, which accounts for means separation at the highest rates tested (Fig. 3, Supplementary Table S1). The disease control effect of the two fungicides combined together was partially additive.

We evaluated chlorothalonil as an "add-on" to prothioconazole and benzovindiflupyr. Treatments including chlorothalonil reduced disease significantly across rates of prothioconazole plus benzovindiflupyr. The interaction between chlorothalonil and prothioconazole and benzovindiflupyr was statistically significant up to the quadratic term: The effect of these three fungicides were more pronounced at the higher treatment levels (Fig. 3). Treatments at the 100% and 200% rate levels (PBC100 & PBC200) had very low or zero disease, respectively.

The mean yield in the untreated control was 67 dt ha\(^{-1}\). There was a statistically significant linear effect of fungicides to yield (Fig. 4, Supplementary Table S2). Fungicide treatments were generally additive for yield. There were no interactions between prothioconazole and benzovindiflupyr, but there was a significant linear interaction between chlorothalonil and benzovindiflupyr, illustrating the marginally less efficacy of benzovindiflupyr than combined fungicide treatments at higher rates (Fig. 4, Supplementary Table S2).

The prothioconazole and prothioconazole plus benzovindiflupyr treatment showed a statistically significant causal effect. Total yield increases across rates were 4.5 and 4.8 dt ha\(^{-1}\) respectively. The percent yield increase mediated by AUDPC reduction was 85\% \( (p < 0.05) \) for prothioconazole plus benzovindiflupyr treatment. The correlation between yield and AUDPC (Supplementary Fig. S1) shows that prothioconazole has a steeper yield reduction through higher AUDPC than benzovindiflupyr.

The fungicide treatments resulted in different mutation patterns in \( R. collo-cygni \) populations at both experimental sites (Figs. 1B,C & 2B,C). Due to the clear positive correlation between \( Cyp51 \)-I325T
and Cyp51-I328L ($R^2=0.97$) only data from Cyp51-I325T were shown. In Wels, the population in the untreated control plots reached 3.3 times higher Cyp51-I325T frequencies compared to St. Poelten (53.8% and 16.1%; Fig. 1B & 2B). Frequencies in Cyp51-I325T mutations were more pronounced in the prothioconazole and prothioconazole plus benzoindiflupyr treatments compared to the benzovindiflupyr treatment. In St. Poelten (Fig. 1B), a dose-responsive increase of Cyp51-I325T was shown by applying P and PB. Compared to the untreated control, treatment PB100 and PB200 significantly increased Cyp51-I325T frequencies from 16 to 68%. Chlorothalonil in the treatments PBC100 and PBC200 showed a good performance against RLS and with the used method no resistant strains could be detected. The result for the mutation Cyp51-I325T was similar in Wels (Fig. 2B), whereby especially the PB treatments showed no increase of mutation during increased dose rates. Chlorothalonil treatments (PBC) slightly increased the Cyp51-I325T frequencies to ±70% by applying PBC25, PBC50 and PBC100 and significantly decreased to zero in treatment PBC200.
The point mutations \textit{sdhC}-H146R and \textit{sdhC}-H153R were measured on a low level and in different frequencies. To include both mutations in one value, the cumulative frequency (\textit{cum.}) of \textit{sdhC}-H146R and \textit{sdhC}-H153R mutations was calculated and represented as \textit{cum.-sdhC}. In Wels, \textit{cum.-sdhC} in untreated control reached higher frequencies compared to St. Poelten (Figs. 1C & 2C), in detail \textit{cum.-sdhC} in Wels (10.1%) was 3.6 times higher compared to St. Poelten (2.8%). Compared to the untreated control, changes in frequency of \textit{cum.-sdhC} were more or less stable through prothioconazole and heavily increased through high SDHI dosage of benzovindiflupyr (B) or prothioconazole plus benzovindiflupyr (PB). In both locations, treatments B100, B200, PB100 and PB200 significantly increased the \textit{cum.-sdhC} frequencies compared to untreated control. Additionally, significant differences were found between B100 and B200 as well as between PB100 and PB200. In St. Poelten (Fig. 1C), fungicide application with 75 g benzovindiflupyr (B100) or additionally with 150 g prothioconazole (PB100) increased the \textit{cum.-sdhC} frequencies more than sixfold (from 2.8% to over 18%). At twice the amount (treatment B200 and PB200), \textit{cum.-sdhC} populations as a result of different fungicide treatments in Wels (4 plots/treatment). Different letters above the box plots indicate significant differences according to GT2-Hochberg test (p=0.05).
increased between 17 and 20 times (46.4–56.6%). In Wels, under a higher \textit{cum.-sdhC} population in the untreated control, the increase was not that high compared to St. Poelten. Here, the frequencies increased 2 to 3 times after application B100 and PB100 and 4 times after application B200 and PB200, respectively (Fig. 2C). Within the chlorothalonil treatments, the low dose-rates in treatment PBC25 and PBC50 resulted in a stable \textit{cum.-sdhC} and treatment PBC100 or PBC200 decreased \textit{cum.-sdhC}, with no significant differences compare to untreated control.
Frequency and distribution of *Cyp51*- and *sdhC*-resistant *R. collo-cygni* population in Austria—a field monitoring

An intensive monitoring program over the main Austrian barley growing areas was started in 2017 to get an overview about the frequency of the most important *Cyp51* and *sdhC* mutations. Resistance mutations leading to the amino acid alteration *Cyp51*-I325T, *Cyp51*-I328L, *sdhC*-H146R and *sdhC*-H153R have been identified with pyrosequencing and are shown in Fig. 5 and Supplementary Table S3. In total, 58 barley fields were analysed between 2017 and 2019 from different Austrian sites. In 2018, no *Ramularia collo-cygni* DNA was detected in samples from the Mannersdorf and Leobersdorf sites, and in 2019 from Grafenschlag, Harmannsdorf I and Mannersdorf. Leaves collected in Grafenschlag and Harmannsdorf I showed clear symptoms of *P. teres* not RLS. After microscopic examination of leaves from Mannersdorf and Leobersdorf also no typical RLS symptoms were found.

Frequencies of mutations varied among loci (*p* < 0.0001). The frequencies for each mutation were significantly different from each other, expect for *Cyp51*-I325T versus *Cyp51*-I328L. Estimated *Cyp51*-I325T frequencies were 18.8% and was not significantly different from *Cyp51*-I328L (19.7%). The estimated frequency of *sdhC*-H146R was 2.0%. The alteration *sdhC*-H153R was found in lowest frequencies within an estimate of 1.1%. Estimates of mutation frequencies declined year on year; the year effect was noteworthy though not statistically significant (*p* = 0.0571) at this level of replication (5–27 samples per locus year). 2019 showed a lower frequency with an odds ratio of mutation of 0.65 versus other years. The decline of the frequency of each mutation indicates there is no increase and very likely, a decrease of fungicide resistance loci in the populations surveyed (Fig. 6).

Due to the good correlation within *Cyp51* mutations (*R^2^ = 0.933; field monitoring data; data not shown) only data from *Cyp51*-I325T and *cum.-sdhC*
were shown (Fig. 5). At site Goedersdorf (2019), Kleinwilfersdorf (2017) and Wasenbruck (2019) *R. collo-cygni* was detected with qPCR but only wild-type population. In the 3 samples from Gerhaus, Preuwitz and Marchtrenk (2019) no *sdhC* mutations were found, although these samples showed Cyp51-I325T frequencies above 20%. All other samples, except the 5 samples which did not contain *R. collo-cygni* DNA, include Cyp51 as well as *sdhC* mutations. Highest amount of Cyp51-I325T alteration was found in Wels and Rohr/Kremstal (above 47%) in 2018 and in Gerhaus 49.5% and Pottenbrunn 72.3% in 2019. Higher frequencies were more often found in north-west Austria compared to south-east Austria, such as Styria. Nevertheless, in some samples from the south-east, we found also high Cyp51-I325T frequencies: 28% in St. Poelten (2018), 24% in St. Paul (2018), and 20% in Wildon (2019). We also observed differences between years: in the southern Austrian region lower frequencies were found in 2019, in the east (near Vienna) no *R. collo-cygni* was detected in 2018 and in 2019 a sample with 50% of Cyp51-I325T was analyzed. *Cum.-sdhC* frequencies were generally lower compared to Cyp51-I325T and frequencies were additionally lower in 2019 compared to 2018. Upper Austria showed the most cases with *cum.-sdhC* frequencies above 6% in 2018. In 2019 only two locations were tested, whereby one sample showed no mutation and the second sample a low *cum.-sdhC* mutation of 4.2%. In the east of Austria very low frequencies were found, except of one sample in Drassmarkt (2018). In the south there were 3 locations with more than 6% *cum.-sdhC* in 2018 and only one location with more than 4% in 2019.

**Evaluation of alternative fungicide strategies**

In 2019, three field trials were run to evaluate new fungicide strategies to protect barley against RLS and other diseases. In this year, the RLS epidemics were moderate: In the untreated control (treatment 1) mean *Ramularia* disease severity on upper two leaves reached 26% in Marchtrenk, 38% in Kirchdorf am Inn and 41% in St. Poelten (Table 3). With chlorothalonil in combination with prothioconazole plus benzovindiflupyr (treatment 3), RLS severity were reduced to under 2.6%, which leads in an efficacy of over 94%. Prothioconazole plus benzovindiflupyr (treatment 2) resulted in an efficacy between 51 and 90%. Prothioconazole plus benzovindiflupyr showed significantly lower disease severity compared to the untreated control and, in addition, significantly weaker performance than prothioconazole plus benzovindiflupyr plus chlorothalonil in Kirchdorf am Inn and St. Poelten. Adding of sulfur (treatment 4) resulted in a slightly benefit in RLS control compared to prothioconazole plus benzovindiflupyr but was not as good as treatments including folpet. Folpet treatments (treatment 5, 6 and 7) provided an efficacy between 80 and 95%—all three treatments did not differ

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**Fig. 6** Model fit estimated frequencies of tested mutations (*sdhC*-H146R, *sdhC*-H153R, Cyp51-I325T and Cyp51-I391L) over the years. The error bars show the 95% confidence limits (CI)
Table 3 Mean RLS disease severity (DS %) and yield (dt ha\(^{-1}\)) of three field trials in 2019. Treatments within the same category with the same letter are not significantly different (p = 0.05). GT2-Hochberg test. Data represent the mean ± SE.

| Treatment* | Kirchdorf am Inn | Marchtrenk | St. Poelten | Yield*** Increase in dt ha\(^{-1}\) | Yield*** Increase in yield (%) |
|------------|-----------------|------------|-------------|----------------|-----------------------------|
| DS (%)**   | ± SE            | Efficacy (%) | DS (%)** | ± SE | Efficacy (%) | DS (%)** | ± SE | Efficacy (%) | dt ha\(^{-1}\) | ± SE | yield (%) |
| 1          | 37.50           | 1.3 a       | 25.53      | 2.94 a | 41.41 | 3.4 a | 101.31 | 2.2 b | -              |
| 2          | 11.69           | 1.8 b       | 2.44       | 0.15 b | 20.44 | 4.1 b | 110.11 | 2.7 a | 9              |
| 3          | 1.41            | 0.2 d       | 0.78       | 0.21 b | 2.56  | 0.5 d | 112.55 | 2.7 a | 11             |
| 4          | 8.72            | 0.3 bc      | 2.09       | 0.61 b | 14.28 | 3.4 bc | 111.88 | 2.3 a | 10             |
| 5          | 5.56            | 1.1 cd      | 1.34       | 0.35 b | 7.00  | 3.0 cd | 111.95 | 2.6 a | 11             |
| 6          | 5.34            | 0.2 cd      | 1.34       | 0.12 b | 8.13  | 2.2 cd | 114.39 | 2.5 a | 13             |
| 7          | 8.09            | 0.7 bc      | 1.19       | 0.25 b | 6.28  | 1.1 cd | 112.48 | 2.5 a | 11             |
| Treatment (T) | < .001       | -          | < .001     | -     | < .001 | -     | < .001 | -             |
| Location (L) | -             | -          | -          | -     | -     | -     |        | 0.9787       |
| TxL        | -              | -          | -          | -     | -     | -     |        |               |

* according to Table 2
** mean disease severity of upper two leaves
*** average of all sites
significantly. Average yield of all trials in untreated control was 101.31 dt ha$^{-1}$. All fungicide treatments significantly increased the yields.

Discussion

The resistance of fungal pathogens to DMI and SDHI fungicides is an ongoing development. Only by combining all relevant strategies to minimize the population of $R.~collo-cygni$ it will be possible to slow down the selection process. In our dose-rate experiments four rates of fungicides and three active ingredients in four different strategies have been pursued to explain their impact on disease severity, GLA, yield and the alteration of $\text{Cyp51}$ and $\text{sdhC}$ mutations. In 2018, for both sites, a high RLS pressure was observed and resulted in a serious disease severity. Our results indicate that RLS severity was higher for treatments that included a single fungicide (prothioconazole or benzovindiflupyr). AUDPC was reduced by 39% through prothioconazole and 52% through benzovindiflupyr, both measured at double the recommended dose rate (P200 and B200). The effects on RLS AUDPC of prothioconazole and benzovindiflupyr were partially additive across rates, and the combination outperformed prothioconazole alone for AUDPC across all rates. There was a tendency for slightly less than expected benzovindiflupyr efficacy at higher rates of prothioconazole. This underlines the principal that two agents with different modes of action should perform additively until dose rate sufficiency is achieved. The mix including chlorothalonil showed much lower AUDPC across rates than that without, including almost no disease at the highest rates. Hence, it was not possible to achieve optimal disease control in this study without the previously most important but now banned active ingredient, chlorothalonil.

Our data from fields with DMI and SDHI resistant $R.~collo-cygni$ populations, indicate single use of prothioconazole and benzovindiflupyr is not enough to achieve high yields. Additionally, treatments containing multiple chemistries including the multi-site inhibitor chlorothalonil were consistently associated with the highest yields. However, at this level of replication (two sites), the effect of PB was statistically indistinguishable from PBC for yield, even though the effect of PBC on AUDPC was much higher than PB. Hence, this study suggests that the combination of P and B together are necessary to protect yield in the absence of the full disease control conferred by chlorothalonil.

The differences between treatments in the resulting yields were not as pronounced as for disease severity given this level of replication. A recent study also reported that the significant impact of fungicides is not necessarily reflected to the same increase in yield (Stetkiewicz et al., 2019). They pointed out that soil type and soil quality has also a high impact and moreover, that the differences in yield were lower in drier years. Yield is a response to many more factors than disease responses to fungicides. As for many other pathogens, precipitation and temperature are significant factors in yield and disease severity (Wiik & Ewaldz, 2009). Our experiments were conducted under drier and warmer conditions compared to the long-term average, especially the dry weather conditions before harvest resulted in a rapid ripening and this may have resulted in lower effect of fungicide applications on yield increase. Similarly, a study in wheat and barley in Nordic-Baltic region showed that biotic and abiotic stress like drought affected yield (Jalli et al., 2020). In addition, with other stress factors this could also resulted in decreased yields. Nevertheless, the results of this study showed a causal effect of P and PB treatments for yield, with 85% of the yield increases in the PB treated plots attributable to disease suppression. Altogether, this suggests that PB can protect yields adequately, even in the absence of complete disease control.

The steeper AUDPC reduction at lower prothioconazole or benzovindiflupyr rates in St. Poelten than Wels suggests that there is a higher threshold of these two active ingredients for control at site Wels, where frequencies of resistant population is higher and more developed. This likely reflects that the low to moderate frequencies of $\text{Cyp51}$ and $\text{sdhC}$ mutations are the main reasons behind low disease decrease for these fungicide groups. Recurrent treatments with the same fungicide can lead to an increase of resistant mutations in pathogen populations. If then the frequency of mutation becomes a considerable proportion of the population, the efficacy of fungicides can be affected (Torriani et al., 2017). Alternatively, a fungicide strategy which include more active ingredients leads to better disease control, previously reported for pathogens of wheat (Heick et al., 2017a; Lynch et al., 2017; Sylvester & Kleczewski, 2018; Wieczorek et al.,...
Interestingly, chlorothalonil was associated with greater disease reduction at higher Cyp51 and cum.-sdhC frequencies in the untreated control at our Wels experimental site than at St. Poelten, especially at lower prothioconazole and benzovindiflupyr rates. This underlines the power of the multi-site inhibitor in a R. collo-cygni population with higher frequencies of tested mutations and supports the importance of rotating modes of action and combining different modes of action to prevent fungicide resistance.

Higher yields were connected to lower RLS disease severity and greener leaves. The AUGLAC is affected by diseases and leaf senescence. The effect of fungicides on leaf senescence is complex and may also be related to the host physiology (Cromey et al., 2004). However, in our study leaf senescence was mainly determined by a short hot period before harvest.

There are several studies on DMI and SDHI resistance of fungal species from field and laboratory mutants of fungi (Avenot & Michailides, 2010; Cools, 2008; Heick et al., 2017a; Piotrowska et al., 2019; Rehfus et al., 2019; Torriani et al., 2017; Wieczorek et al., 2015), but there are no previous field studies on the effect of fungicide dose-rates on the quantitative selection of Cyp51 and sdhC mutations on R. collo-cygni populations. In our study, we found that higher dose-rates of prothioconazole lead in higher Cyp51 frequencies and higher dose-rates of benzovindiflupyr lead in higher sdhC frequencies. A selection for fungicides is described for other pathogens and same or other single-site fungicides (Dooley et al., 2016; Heick et al., 2017a; Simoes et al., 2018; Van den Bosch et al., 2011; Zhou & Mehl, 2020). The review by Van den Bosch et al. (2011) mentioned that field experiments are difficult to determine the effect of fungicide doses on fungicide resistance, except for the selection phase in resistance development. However, they documented that the selection of resistant types mostly increases with higher dose rates. These findings are in line with data of quantitative detection of Cyp51-I325T and cum.-sdhC mutations in this study. Cyp51-I325T frequencies increased with increasing prothioconazole rates and reached a frequency of over 95% for site Wels. This means that with an application of 300 g ha⁻¹ prothioconazole (P 200) almost the whole R. collo-cygni population is less sensitive to this DMI. Based on a lower Cyp51-I325T frequency in untreated control in St. Poelten, an application with 300 g ha⁻¹ prothioconazole (P 200) increased the Cyp51-I325T frequency to over 60%, which means a large increase of 3.8 times. The study by Heick et al. (2017a) reported changes in Cyp51 frequencies after fungicide application in Z. tritici population. They noted a marked increase after three applications of DMI’s compared to one application with DMI plus SDHI. Mutations in the sdh-C gene resulted in amino acid changes at codon 146 (sdhC-H146R) and 153 (sdhC-H153R), both represented in cum.-sdhC frequencies. These frequencies were also increased through SDHI application. Comparable for Cyp51-I325T mutation, cum.-sdhC showed a major increase in St. Poelten, where a lower cum.-sdhC frequency in untreated control was present, compared to Wels. The same trend was shown for sdhC-I86F in Phakopsora pachyrhizi (Simoes et al., 2018). The authors found in a population with lower sdhC-I86F frequencies a major increase after SDHI application and in populations with moderate to high sdhC-I86F frequencies no or just a small increase after SDHI application. The benzovindiflupyr treatments (B25-B200) also increased Cyp51-I325T frequencies which is not comparable with the prothioconazole treatments (P25-P200) where no increase in cum.-sdhC frequencies were found. However, there is a lack in field research on R. collo-cygni and therefore the reason why one fungicide increases the mutation from the other fungicide group is not easily to justify. Several conclusions could be drawn from studies on Z. tritici, which is closely related to R. collo-cygni (McGrann et al., 2016; Stam et al., 2018). The population of Z. tritici is very heterogeneous in a single field or single trial (Stammler & Semar, 2011)–this can lead to different DMI sensitivities and Cyp51 frequencies within one field or treatment. Moreover, earlier studies reported that different DMIs have a different impact on selecting for Cyp51 mutations (Cools, 2008; Heick et al., 2017a; Wieczorek et al., 2015). The most plausible explanation is provided by Stracke et al. (2018), who showed that R. collo-cygni isolates pretreated with prothioconazole were less sensitive to prothioconazole and additionally to benzovindiflupyr compared to the untreated control. Furthermore, the study by Rehfus et al. (2019) came to the result that isolates with low levels of DMI adaptation were connected
to mutation in the SDH gene. This strong connection could lead to an increase of Cyp51 mutations after SDHI application. To elucidate why an SDHI application is increasing Cyp51 mutations, further work is needed. However, chlorothalonil in higher rates could hold Cyp51-I325T and sdhC mutations at lower levels.

SDHIs and DMIs are key fungicides for disease management in barley and therefore the Fungicide Resistance Action Committee (FRAC) has been conducting monitoring programs in Europe for many years. In 2016, the first shifted or resistant isolates were found in Upper- and Lower Austria, especially in samples collected after SDHI application (Torriani S. and Bernhard-Frey R. 2019, personal communication). Our study is the first extensive monitoring of fungicide resistance within R. collo-cygni populations in Austria and the results are important for further management strategies in barley. Out of 58 samples in 2017, 2018 and 2019, only in five samples R. collo-cygni could not be detected. This shows that R. collo-cygni is present in all main barley growing areas, the Alpine-north climate region, the Pannonian climate region, the Illyrian and the Alpine-south climate region. In the Pannonian climate region, characterized through dry spring and summer periods, especially in the region east from Vienna, no Ramularia collo-cygni was detected in 2018, but was found in 2019. This is contrary to the first Austrian findings of Huss et al. (2009) which refers to the lack of RLS in the dry eastern part of Austria. Hoheneder et al. (2021) showed that RLS level, sporulation rate and DNA content varied between years or different weather conditions. Especially long drought periods suppress RLS under field conditions. Our visible observations over several trials in different locations over the last years in Austria arrived at the same conclusions. R. collo-cygni with its large population and high mutation rate can therefore easily adapt to environmental conditions that may be exacerbated with climate change, which is increasing the range of the pathogen (Piotrowska et al., 2016; West et al., 2012).

The mutation development to DMI and SDHI fungicides in Western Europe is an ongoing process, especially the combined resistance towards SDHIs and DMIs (ReHFus et al., 2019). This is in line with findings in our study. In almost all cases Cyp51-I325T were detected together with sdhC mutations. Cyp51-I325T mutation were found in higher frequencies compared to cum.-sdhC mutation – resulting in a lower performance of DMIs. Especially for Cyp51-I325T mutation, we found higher frequencies in the north compared to the south-east of Austria. This indicates that the resistance development has started in north-west and a north-west to south-east gradient exists. The moderate to high frequencies in Germany and no to low frequencies in Slovenia and Croatia as well as no Cyp51 mutation in Italy (FRAC 2020) support this assumption. Based on our study, it can be concluded that resistant strains of R. collo-cygni population did not increase over the last few years due to the well-established process of chlorothalonil application in Austria since the first detection of resistance. The frequency of Cyp51 and sdhC mutations was significantly lower in 2019 compared to the previous two years. This decrease is associated with the general lower disease pressure in 2019 and the wide use of chlorothalonil. The lower frequency of mutations shows that DMIs and SDHIs still have performance in RLS control and mainly increase in yield is given through use of these fungicides. Nevertheless, an intensive and carefully considered management strategy is important to prolong the lifespan of DMIs and SDHIs.

The control on RLS heavily depended on the use of chlorothalonil; however, this multisite fungicide will not be available in future EU fungicide strategies due to recent regulations. Thus, QoIs, DMIs and SDHIs are brought more into focus, but in case of R. collo-cygni control all three groups are affected by site mutations, resulting in shifts in efficacy. The adding of sulphur, azoxystrobin and especially folpet in fungicide strategies with DMI plus SDHI resulted in a better RLS control compared to DMI plus SDHI alone. This shows that a fungicide strategy including more active ingredients from different fungicide groups are responsible for a better RLS control. Fungicide mixtures are also the key to slow down the emergence and selection of fungicide mutations and prolong effective life of fungicides (Van den Bosch et al., 2011). The loss of chlorothalonil will rapidly drive the spread and increase of resistance (Kildea et al., 2018). To counteract this, the authors refer to a moderate control where folpet is used and outlined the loss of around 10% in yield through the replacement of chlorothalonil with folpet. The improved performance of folpet compared to sulphur were also raised by Stracke et al. (2018). The adding of azoxystrobin in the tank mix with prothioconazole plus benzovindiflupyr
plus folpet showed no benefit in RLS control, but increased yield by 2%. The G143 point mutation as described in Fountaine and Fraaije (2009) is responsible for the missing effectiveness of azoxystrobin leads in a delay senescence and as a result increase in yield (Jorgensen & Olesen, 2002). The disease severity in this trial was not very high, but folpet looks like a good replacement. Further investigations are necessary, especially to show if the selection process for DMIs and SDHI mutations can be slow down and delay to protect DMIs and SDHIs in their efficacy.

Barley growers should adjust to the fact that RLS is present in the main Austrian barley growing areas and disease level will increase due to good adaptability to climate change. Based on the low to moderate Cyp5f1 and sdhC frequencies and as well to protect other barley diseases, DMIs and SDHIs are essential to protect barley. To improve the lifetime of compounds with these modes of action, a focused application between GS 49 and 59 associated with the lowest recommended dose-rates of DMIs and SDHIs in combination with other active ingredients like folpet are essential, even if the combination of prothioconazole and benzovindiflupyr protected the yield under current situation. In the absence of new fungicides against RLS control, current fungicides have to be carefully used to prolong their activity as long as possible. From the agronomic point of view, this is a compromise to ensure yield and quality in barley.

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Declarations

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