Effect of year, sampling month and grape cultivar on noble rot incidence, mycelial growth rate and morphological type of *Botrytis cinerea* during noble rot development

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**Abstract** The aims of this three-year study were firstly to investigate the effect of 3 years, three sampling months and two grape cultivars (cvs) on noble rot incidence in the field and on the mycelial growth rate of *Botrytis cinerea* isolates in vitro under three incubation temperatures, and secondly to show possible effects of years, sampling months and cultivars on eight morphological (four mycelial: M I-IV and four sclerotial: S I-IV) types of *B. cinerea* isolates incubated at 20 °C. In addition, the relationship between monthly noble rot incidences and morphological types was calculated. Analyses of variance indicated significant differences among years, months and cultivars for noble rot incidence. Noble rot incidences were significantly higher on cv. ‘Turán’ compared to cv. ‘Olaszrizling’ in November in all years. Noble rot incidences were significantly lower in September compared to values in November in all years. Analyses of variance indicated significant differences among years, months, and temperatures for mycelial growth rate but the differences among cultivars were non-significant. Mycelial growth rates at 15 and 25 °C were higher in 2013 compared to 2014 and 2015; however, the rates at 20 °C were higher in 2014 compared to 2013 and 2015. The overall mean growth rate was the highest in 2013 (76 mm) compared to either 2014 or 2015 (65 or 69 mm, respectively). Mycelial growth rates were significantly different between earlier (September) and later (November) sampling months in three cases: i) in 2013 at 15 °C, ii) in 2014 at 20 °C, and iii) in 2015 at 25 °C. Analyses of variance indicated significant differences among years and months for the frequency distribution of the morphological types but the differences among cultivars were non-significant. In general, the most frequent mycelial and sclerotial types were M I-III and S IV (means were 21.5, 16.1, 28.9, and 13.1%, respectively). Isolates of mycelial types were more frequent in 2013 (Σ M-type 82.2%) compared to 2015 (Σ M-type 57.1%). The frequency distributions of morphological types among sampling months showed differences between the earlier (September or/and October) and the later months (November) in 2013 for M I-III, S II and S IV types; in 2014 for M I, M III, M IV, S I, S III and S IV types; and in 2015 for M I, M IV and S I-IV types. Pearson correlation analyses revealed a
Botrytis cinerea is a common and widespread ascomycete fungus and necrotrophic pathogen attacking numerous different plant species e.g. grape, tomato, kiwi fruit, strawberry and raspberry, herbaceous, shrub and tree species (Holz et al. 2007). In grape, the pathogen can cause gray mold resulting rot of berries but under certain microclimatic conditions, infected berries go through the process of noble rot (Magyar 2011; Fournier et al. 2013).

Botrytis cinerea populations were widely studied in European wine regions suggesting that several factors can influence the population structure such as host, geographic origin, mode of reproduction, occurrence of mutations, the presence of transposon elements, and/or gene-flow (Campia et al. 2016). Geographical diversity among the pathogen populations was associated with gray mold versus noble rot symptoms in various wine regions of France (Fournier et al. 2013; Walker et al. 2015). Authors showed that noble rot development occurred under the combinatory effects of climate, cultivar, year, cultural practices, and berry microbiome (Ribéreau-Gayon et al. 2006; Blanco-Ulate et al. 2015). The occurrence of noble rot is dependent mainly on microclimatic conditions (Ciliberti et al. 2015b; Elad et al. 2016); however, there is a lack of information that how the different microclimatic conditions are connected to B. cinerea phenotypes (Martinez et al. 2003; Martinez et al. 2005; Váczy et al. 2008; Ciliberti et al. 2015a; Ciliberti et al. 2015b) but isolate features were not investigated in relation to the early or late autumn period of noble rot development on various cultivars.

Mycelial growth rate, as one of the basic isolate features, was studied widely for noble rot caused by B. cinerea (Youssef and Roberto 2014). An intensive mycelial growth resulted in a higher infective capacity, and the lower infective capacity was connected to a less sclerotia production (Cantoral et al. 2011; Vallejo et al. 2003). A more wider phenotypic characterizations of B. cinerea strains during noble rot were performed for instance by describing the morphological types and features (e.g. Martinez et al. 2003; Lorenzini and Zapparoli 2014) or the effect of water activity on mycelial and conidial growth rates of the B. cinerea strains (Rousseau and Doneche 2001). However, the connections among the morphological types, sampling months or grape cultivars were not investigated on isolates originated from various periods of noble rot development.

The aims of this three-years study were: firstly, to investigate the effect of 3 years, three sampling months during noble rot development and two grape cultivars on noble rot incidence in the field and on the mycelial
growth rate of *B. cinerea* isolates under three incubation temperatures; secondly, to show possible effects of years, sampling months and cultivars on eight morphological (four mycelial and four sclerotal) types of *B. cinerea* isolates incubated at 20 °C; and thirdly, to calculate the overall relationship between monthly noble rot incidences and morphological types.

The study was performed in three consecutive years (2013–2015) in a vineyard located at Somolya (47° 54' 00" N; 20° 30' 00" E) in the Eger wine region, Eastern-Hungary. The main soil type was brownish forest soil with riolit tufa stones. The vineyard was planted in 1998 with planting distance between rows of 2.9 m and within rows of 0.9 m. A guyot pruning type has been performed since the establishment of the vineyard. Two cultivars were grown: the white-skinned ‘Olaszrizling’ and the red-skinned ‘Turán’. Cultivar (cv) ‘Olaszrizling’ is declared as suitable for botrytised wine making (Robinson et al. 2013) and cv ‘Turán’ can be used for a unique red botrytised wine (Robinson et al. 2013). In the vineyard, fungicides were used in all years form early April to mid-July (Table 1). All sprays were applied with a Lochmann RPS10/70 axial airblast sprayer (Lochmann Plantatech, Nalles, Italy) with a ceramic hollow cone at 0.5 to 1.5 MPa with a volume of 1000 l ha⁻¹. Sprays against insects were not applied in either years and a mechanical weed management was applied two to three times annually in the vineyard.

Rainfall (mm) and daily temperature (mean, minimum and maximum; °C) were detected using a Boreas agrometeorological station (Boreas Ltd., Érd, Hungary) in the vineyard from 1 September until 30 November in the 3 years.

Three sampling months (September, October and November) were chosen according to the 3-month-period of noble rot development. According to this, noble rot incidence was assessed in 12 September, 27 October and 28 November in each year. Grape clusters of 20 plants were assessed on each cultivar in each sampling date. A grape bunch was considered to be noble rotted if at least one berry showed the typical symptoms caused by *B. cinerea*. Noble rot incidence was calculated for each observed plant. Grape berries with symptoms of noble rot were collected from the vineyard at the same dates in each year. A minimum of 20 samples of naturally-botrytised grape berries was collected in each month-year combination and on each cultivar. Then *B. cinerea* from the samples were cultured on rose bengal chloramphenicol (RBC) medium (Scharlab S.L., Spain). Then single-spore cultures were selected and maintained on potato dextrose agar (PDA) (Biolab Zrt, Budapest, Hungary). For growth tests, a 5-mm mycelial plug was cut from the margin of an actively growing 4-day-old culture and placed in the center of a Petri dish containing PDA. Plates were incubated at 20 °C (optimum temperature) and at 15 and 25 °C (sub-optimum temperatures) in dark. Mycelial growth diameter was measured to a 4-day post inoculation. Three replicates were performed for each isolate and the experiment was carried out twice.

Morphological features of the same 20 isolates were determined for the 3 years, three sampled months and the two cultivars in order to investigate their effects on the morphological types of *B. cinerea* isolates. Morphological types were characterized after a 21-days inoculation but only on those isolates that were grown at the optimum temperature (20 °C). Then cultures were macroscopically examined for sporulation, mycelium and sclerotium production (Table 3). Then isolates were classified into the morphological groups of mycelial (M) or sclerotial (S) types, which contained four mycelial (M I, M II, M III, M IV) and four sclerotal (S I, S II, S III, S IV) types according to Martinez et al. (2003) (Table 2). Three replicates were performed for each isolate and the experiment was carried out twice.

Data on noble rot incidence were analysed by using analyses of variance (ANOVA) in order to determine the effect of year, sampling month, cultivar, and their interactions. Data on mycelial growth rate were also analysed by using ANOVA in order to determine the effect of year, sampling month, cultivar, temperature and their interactions. For both measures, means were separated by using an LSD test (α = 0.05). Frequency distribution (FD%) of each morphological type was calculated for each year, sampling month and cultivar as FD% = (isolate numbers in a given classes of morphological type / total numbers of isolates) × 100. Frequency distributions were then analysed by using ANOVA in order to determine the effect of year, sampling month, cultivar, morphological group and their interactions. Means were separated by using an LSD test (α = 0.05). Before the analyses, frequency distributions were arcsine-square root transformed for data normality.

Pearson correlation between morphological types (Σ S-type and Σ M-type) and monthly noble rot incidences (September, October and November) was calculated for the combined data set of the 3 years and two cultivars.
All analyses were done using Genstat 5 Release 4.1 (Lawes Agricultural Trust, IACR, Rothamsted, UK).

The September daily temperatures were the highest in 2015, the October ones in 2013, while the November ones in 2014 (Table 3). During the three autumn months, the maximum values of daily temperature were 22.7, 23.1, and 24.0 °C in 2013, 2014, and 2015, respectively. The total amount of rainfall was the lowest (112.5 mm) in the autumn of 2013 and the highest (199.4 mm) in the autumn of 2015 (Table 3). Weather conditions for noble rot development were more favourable in the autumn of 2013 compared to either the autumn of 2014 or 2015.

Analyses of variance indicated significant differences among years, months, and cultivars (mean square = 874.4, 14,201.2, and 2035.4; \( P < 0.001 \), <0.001, and <0.001, respectively) for noble rot incidence. In addition, all interactions among treatment factors were non-significant. Across the 3 years and two cultivars, noble

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### Table 1 Fungicide spray schedules from early April to mid-July in the experimental vineyard (2013–2015, Somolya, Hungary)

| Date       | Active ingredients                        | Trade name\(^b\) | Dosage (%) |
|------------|-------------------------------------------|------------------|------------|
| 2013       |                                           |                  |            |
| 10 April   | copper hydroxide, 24%; plant oil, 25%     | Vegesol RS       | 0.4        |
| 1, 10 May  | elemental sulphur, 80%                    | Kumulus S        | 0.4        |
|            | mancozeb, 750 g kg\(^{-1}\)              | Vondozeb DG      | 0.2        |
| 24 May, 11 June | elemental sulphur, 80%              | Kumulus S        | 0.4        |
|            | dimetomorf, 9%; mancozeb, 60%           | Acrobat MZ WG    | 0.2        |
| 25 June, 5 July | elemental sulphur, 80%                | Kumulus S        | 0.4        |
|            | copper hydroxide, 77%                    | Champion WG      | 0.2        |
| 15 July    | elemental sulphur, 80%                   | Kumulus S        | 0.4        |
| 2014       |                                           |                  |            |
| 12 April   | copper hydroxide, 24%; plant oil, 25%     | Vegesol RS       | 0.4        |
| 5, 16 May  | elemental sulphur, 80%                    | Kumulus S        | 0.4        |
|            | mancozeb, 750 g kg\(^{-1}\)              | Vondozeb DG      | 0.2        |
| 25 May, 13 June | elemental sulphur, 80%                | Kumulus S        | 0.4        |
|            | dimetomorf, 9%; mancozeb, 60%           | Acrobat MZ WG    | 0.4        |
| 24 June, 5 July | elemental sulphur, 80%                | Kumulus S        | 0.4        |
|            | copper hydroxide, 77%                    | Champion WG      | 0.2        |
| 10 July    | elemental sulphur, 80%                   | Kumulus S        | 0.4        |
| 15 July    | elemental sulphur, 80%                   | Kumulus S        | 0.4        |
| 20 July    | meptildinokap, 350 g l\(^{-1}\)          | Karathane Star   | 0.075      |
| 2015       |                                           |                  |            |
| 10 April   | copper hydroxide, 24%; plant oil, 25%     | Vegesol RS       | 0.4        |
| 2, 12 May  | elemental sulphur, 80%                    | Kumulus S        | 0.4        |
|            | mancozeb, 750 g kg\(^{-1}\)              | Vondozeb DG      | 0.2        |
| 22 May, 9 June | elemental sulphur, 80%                | Kumulus S        | 0.4        |
|            | dimetomorf, 9%; mancozeb, 60%           | Acrobat MZ WG    | 0.2        |
| 23 June, 4 July | elemental sulphur, 80%                | Kumulus S        | 0.4        |
|            | copper hydroxide, 77%                    | Champion WG      | 0.2        |
| 15 July    | elemental sulphur, 80%                   | Kumulus S        | 0.4        |

\(^a\) Sprays against insects were not applied in either years and a mechanical weed management was applied two to three times annually in the vineyard

\(^b\) Vegesol RS (BVN Ltd, Budapest Hungary), Kumulus S (BASF Hungary Ltd., Budapest, Hungary), Acrobat MZ WG (BASF Hungary Ltd., Budapest, Hungary), Vondozeb DG (UPL Europe Ltd., Cheshire, UK), Champion WG (Kwizda Agro Ltd, Budapest, Hungary), Karathane Star (Dow AgroSciences Hungary Ltd, Budapest, Hungary)
rot incidence ranged from 0.5 to 10.3%, from 4.3 to 39.4%, and from 26.5 to 78.7% for the sampling months of September, October, and November, respectively (Table 4). The highest noble rot incidence (78.7%) was assessed on cv. ‘Turán’ in November 2015 while the lowest one (0.5%) on cv. ‘Olaszrizling’ in September 2013. Noble rot incidences were significantly higher on cv. ‘Turán’ compared to cv. ‘Olaszrizling’ in November in all years. Noble rot incidences were significantly lower in September compared to values in November in all the 3 years (Table 4).

Analyses of variance indicated significant differences among years, months, and temperatures (mean square = 13,192.4, 4448.9, and 236.5; \( P < 0.001, =0.037, \text{ and } <0.001, \text{ respectively} \) for mycelial growth rate but the

### Table 2 Morphological types of *Botrytis cinerea* isolates grown on PDA medium at 20 °C according to Martinez et al. (2003)

| Morphological type | Mycelium | Sporulation | Sclerotia |
|--------------------|----------|-------------|----------|
| type               | M I      | M II        | M III    |
|                    | Short    | Aerial      | Aerial   |
|                    |          |             |          |
|                    |          |             |          |
| Mycelial masses    |          |             |          |
| Sporulation\(a\)  | −        | +           | ±        |
| Sclerotia\(b\)    | −        | 0           | −        |

### Table 3 Daily temperatures (mean, minimum and maximum) and rainfall in September, October and November in the experimental vineyard (2013–2015, Somolya, Hungary)

| Year and Month | Daily temperature (°C) | Rainfall (mm) |
|---------------|------------------------|---------------|
|               | Mean | Minimum | Maximum | Mean | Minimum | Maximum |
| 2013          |      |         |         |      |         |         |
| September     | 15.4 | 5.8     | 28.0    | 27.4 |         |         |
| October       | 13.3 | 0.6     | 24.6    | 40.5 |         |         |
| November      | 8.1  | −4.0    | 15.5    | 44.6 |         |         |
| Overall autumn\(a\) | 12.3 | 0.8     | 22.7    | 112.5 |         |         |
| 2014          |      |         |         |      |         |         |
| September     | 17.9 | 6.8     | 28.4    | 91.8 |         |         |
| October       | 12.5 | 2.6     | 23.7    | 50.0 |         |         |
| November      | 7.8  | −1.9    | 17.2    | 11.6 |         |         |
| Overall autumn\(a\) | 12.7 | 2.5     | 23.1    | 153.4 |         |         |
| 2015          |      |         |         |      |         |         |
| September     | 18.7 | 9.2     | 34.8    | 54.8 |         |         |
| October       | 11.1 | 3.2     | 23.3    | 122.0|         |         |
| November      | 7.2  | −2.2    | 13.8    | 22.6 |         |         |
| Overall autumn\(a\) | 12.3 | 3.4     | 24.0    | 199.4 |         |         |

\(a\) Average values of the three autumn months for each daily temperature parameter (mean, minimum and maximum) and sum of rainfall in the three autumn months

### Table 4 Noble rot incidence (%) caused by *Botrytis cinerea* on grape clusters of cultivars ‘Turán’ and ‘Olaszrizling’ in September, October and November in the experimental vineyard (2013–2015, Somolya, Hungary)

| Year and Month | Cultivar       | LSD\(_{0.05}\) | Overall |
|---------------|----------------|---------------|---------|
|               | ‘Turán’        | ‘Olaszrizling’ |         |
| 2013          |                |               |         |
| September     | 5.1 a\(b\)    | 0.5 aB        | 3.7     | 2.6 a |
| October       | 7.1 a          | 4.3 a         | ns\(c\) | 5.7 a |
| November      | 50.6 bA        | 28.4 bB       | 7.6     | 39.5 b |
| LSD\(_{0.05}\) | 7.0            | 4.2           |         | 5.6   |
| 2014          |                |               |         |
| September     | 10.3 a         | 7.2 a         | ns\(c\) | 8.8 a |
| October       | 39.4 bA        | 24.3 bB       | 6.9     | 31.9 b |
| November      | 37.6 bA        | 26.5 bB       | 6.6     | 32.1 b |
| LSD\(_{0.05}\) | 6.9            | 5.8           |         | 6.7   |
| 2015          |                |               |         |
| September     | 4.2 a          | 0.8 a         | ns\(c\) | 2.5 a |
| October       | 9.7 a          | 7.3 a         | ns\(c\) | 8.5 a |
| November      | 78.7 bA        | 57.6 bB       | 8.9     | 68.2 b |
| LSD\(_{0.05}\) | 11.5           | 7.4           |         | 9.5   |

\(a\) LSD\(_{0.05}\) = Least Significant Difference at \( P = 0.05 \)

\(b\) For each column, different small letters indicate significant differences at \( P = 0.05 \) among years, months. For each row, different capital letters indicate significant differences at \( P = 0.05 \) among cultivars within a given month

\(c\) ns = non-significant
differences among cultivars were non-significant (mean square = 138.7, $P = 0.164$). In addition, all interactions among treatment factors were non-significant. As cultivar being non-significant, data of mycelial growth rate were averaged for the two cultivars and not shown separately for temperatures, years and months (Table 5). Across the 3 years and the three sampling months, mycelial growth rate ranged from 59 to 79 mm, from 75 to 84 mm, and from 55 to 77 mm for incubation temperatures of 15, 20, and 25 °C, respectively (Table 5). The highest mycelial growth rate (84 mm) was measured for the incubation temperature of 20 °C on isolates collected in September 2014 while the lowest one (55 mm) for the incubation temperature of 25 °C on isolates collected in September 2014. Mycelial growth rates at 15 and 25 °C were higher in 2013 compared to 2014 and 2015; however, the growth rates at 20 °C were higher in 2014 compared to 2013 and 2015 (Table 5).

Table 5 Mycelial growth rate (mm) of *Botrytis cinerea* after 4 days incubation at 15, 20, and 25 °C in 3 years (2013, 2014, and 2015) during 3 months (September, October, and November) of noble rot development (Somolya, Hungary)

| Year, Month  | Temperature range | LSD$_{0.05}$ $^a$ | Overall Temp $^b$ |
|--------------|-------------------|-------------------|-------------------|
|              | 15 °C  | 20 °C  | 25 °C  |                |                |
| 2013         |        |        |        |                |                |
| September    | 79 bB  | 79 B   | 74 A   | 3.6            | 77              |
| October      | 76 ab  | 76     | 74     | ns $^d$        | 76              |
| November     | 75 a   | 77     | 77     | ns             | 76              |
| LSD$_{0.05}$ | 3.4    |        |        |                | ns              |
| 2014         |        |        |        |                |                |
| September    | 59 A   | 84 bB  | 55 A   | 7.5            | 66              |
| October      | 59 A   | 83 bB  | 56 A   | 7.4            | 66              |
| November     | 61 A   | 75 aB  | 56 A   | 6.8            | 64              |
| LSD$_{0.05}$ | ns     |        |        |                | ns              |
| 2015         |        |        |        |                |                |
| September    | 61 A   | 75 B   | 67 aA  | 6.9            | 68              |
| October      | 62 A   | 76 B   | 71 aB  | 7.0            | 69              |
| November     | 61 A   | 75 B   | 72 bB  | 7.3            | 69              |
| LSD$_{0.05}$ | ns     |        |        |                | ns              |
| Overall month|        |        |        |                |                |
| September    | 66 A   | 79 B   | 65 A   | 6.5            | 70              |
| October      | 66 A   | 78 B   | 67 A   | 7.4            | 71              |
| November     | 65 A   | 76 B   | 68 A   | 7.1            | 70              |
| LSD$_{0.05}$ | ns     |        |        |                | ns              |
| Overall year |        |        |        |                |                |
| 2013         | 77 b   | 77 ab  | 75 b   | ns             | 76 b            |
| 2014         | 60 aA  | 81 bB  | 56 aA  | 7.9            | 65 a            |
| 2015         | 61 aA  | 75 aB  | 70 bB  | 7.6            | 69 a            |
| LSD$_{0.05}$ | 6.6    | 5.8    | 7.3    |                | 6.4             |
| Overall      |        |        |        |                |                |
| Year and Month| 66 A  | 78 B   | 67 A   | 7.9            | 70              |

$^a$ LSD$_{0.05}$ = Least Significant Difference at $P = 0.05$

$^b$ Overall Temp = overall temperature from the temperature ranges of 15, 20, and 25 °C

$^c$ For each column, different small letters indicate significant differences at $P=0.05$ among years, months. For each row, different capital letters indicate significant differences at $P=0.05$ among temperature ranges

$^d$ ns = non-significant
The overall mean growth rate was the highest in 2013 (76 mm) compared to either 2014 or 2015 (65 or 69 mm, respectively). Mycelial growth rates were significantly different between earlier (September) and later (November) sampling months in three cases i) in 2013 at 15 °C, ii) in 2014 at 20 °C, and iii) in 2015 at 25 °C (Table 5).

Analyses of variance indicated significant differences among years, months, and morphological groups (mean square = 1095.7, 998.6, and 5652.9; \( P = 0.005, =0.006, \) and <0.001, respectively) for the frequency distribution of the morphological types but the differences among cultivars were non-significant (mean square = 127.6, \( P = 0.438 \)). In addition, all interactions among treatment factors were non-significant. As cultivar being non-significant, data on the frequency distribution of the morphological types were averaged for the two cultivars and not shown separately for years, months, and morphological types (Table 6). Frequency distribution of the morphological group of mycelial types (\( \sum M \)-type, ranged from 51.4 to 91.7%, mean 70.5%) were significantly higher compared to the sclerotial types (\( \sum S \)-type, ranged from 8.3 to 48.6%, mean 29.5%) across the 3 years and the three sampling months (Table 6). In general, the most frequent mycelial and sclerotial types were M I-III and S IV (means were 21.5, 16.1, 28.9, and 13.1%, respectively). The highest frequency distribution (58.3%) was measured for M III type on isolates collected in September 2013. Isolates of mycelial types were more frequent in 2013 (\( \sum M \)-type 82.2%) compared to 2015 (\( \sum M \)-type 57.1%). The frequency distributions of morphological types among sampling months showed differences between the earlier (September or/and October) and the later months (November) in 2013 for M I-III, S II and S IV types; in 2014 for M I, M III, M IV, S I, S III and S IV types; and in 2015 for M I, M IV and S I-IV types (Table 6).

Pearson correlation analyses revealed a significant positive relationship between morphological types (\( \sum S \)-type and \( \sum M \)-type) and monthly noble rot incidences (\( r = 0.676 \) and \( P = 0.048 \)). The relationship indicated that the increasing noble rot incidence from September to November resulted in an increasing frequency of the morphological type S while the frequency of morphological type M decreased.

In this study, the thermic optimum was the same for mycelial growth of \( B. \) cinerea strains originated either from red- or white-skinned grape cultivars (non-significant cultivar effect in ANOVA). The greatest growth rate was observed at 20 °C while the growth rate decreased at 15 and 25 °C (Table 5). Our data confirmed most previously reported results, for instance, Martinez et al. (2003) and Ciliberti et al. (2015b) showed that strains of \( B. \) cinerea were similarly affected by temperature, and the greatest mycelial growth rate or infection was observed at 20 °C, while they decreased in the order at 25, 15, and 5 °C. Other previous studies confirmed a same temperature effect on the infection of grape berries by \( B. \) cinerea (Nair and Allen 1993) and on the development of aerial mycelium of \( B. \) cinerea on grape berries (Thomas et al. 1988). Our results on temperature also confirmed the model developed for predicting the combined effect of temperature and the mycelial growth rate of \( B. \) cinerea (Judet-Correia et al. 2010) and the model describing the relationship between the incidence of berry infection and temperature (Broome et al. 1995).

In addition, a mechanistic model of \( B. \) cinerea on grapevines including weather, vine growth stage, and the main infection pathways was developed by González-Domínguez et al. (2015), which also supports the findings in this study. In connection with these infection pathways, the main four grape growth stages (end of flowering, pre-bunch closure, veraison and before harvest) can differ in the tissue susceptibility for the infection. The improved predicting model by González-Domínguez et al. (2015) verified that the early season infection influenced the severity of \( Botrytis \) infection during berry ripening. This indicates that latent infection under favourable conditions could also be a driving factor in this Hungarian study for the noble rot development.

In relation to temperature effect on mycelial growth rate, the year influence on mycelial growth rate was analysed by Damialis et al. (2015). Authors confirmed a significant connection between mycelial growth rate of \( B. \) cinerea and average decadal air temperature from the 1980s to the 2000s. Their findings may support our results on the growth rate differences among years. In 2013, the mean maximum daily temperature during the autumn (22.7 °C) was the closest to the optimum (20 °C) for the greatest mycelial growth of the \( B. \) cinerea. Meanwhile in 2014 and 2015, the differences between the optimum temperature (20 °C) and the mean maximum daily temperatures during the autumn (23.1 and 24.0 °C, respectively) were higher than in 2013, which is in line with a lower mycelial growth rate and a lower overall frequency of M-types in the isolates of 2014 and 2015.
Table 6 Frequency distribution (%) of *Botrytis cinerea* isolates in eight classes of morphological types (M I-IV and S I-IV), 3 years (2013, 2014, and 2015) during 3 months (September, October, and November) of noble rot development (Somolya, Hungary)

| Mycelial (M) type | Sclerotial (S) type |
|-------------------|---------------------|
| M I               | S I                 |
| M II              | S II                |
| M III             | S III               |
| M IV              | S IV                |
| LSD$_{0.05}$      | Σ M-type$^b$        |
| Σ S-type$^b$      |                     |

| 2013               |                     |
| September          |                     |
| M I               | 8.3 aA$^c$          |
| M II              | 25.0 bB             |
| M III             | 58.3 cC             |
| M IV              | 0.0 aA              |
| LSD$_{0.05}$      | 11.7 ns             |
| Σ M-type$^b$      | 16.1 ns             |
| Σ S-type$^b$      | 6.8 ns              |
| October            |                     |
| M I               | 12.5 aAB            |
| M II              | 20.0 abB            |
| M III             | 42.5 bC             |
| M IV              | 0.0 aA              |
| LSD$_{0.05}$      | 15.2 ns             |
| Σ M-type$^b$      | 13.8 ns             |
| Σ S-type$^b$      | 9.9                 |
| November           |                     |
| M I               | 42.2 bC             |
| M II              | 11.1 aA             |
| M III             | 24.4 aB             |
| M IV              | 2.2 A               |
| LSD$_{0.05}$      | 15.2 ns             |
| Σ M-type$^b$      | 12.9 ns             |
| Σ S-type$^b$      | 80.0 ab             |

| 2014               |                     |
| September          |                     |
| M I               | 37.6 bD             |
| M II              | 12.8 B              |
| M III             | 25.4 aC             |
| M IV              | 0.0 aA              |
| LSD$_{0.05}$      | 11.2                 |
| Σ M-type$^b$      | 12.1                 |
| Σ S-type$^b$      | 75.0 b              |
| October            |                     |
| M I               | 30.8 bB             |
| M II              | 11.5 A              |
| M III             | 38.5 bB             |
| M IV              | 0.0 aA              |
| LSD$_{0.05}$      | 10.8                 |
| Σ M-type$^b$      | 11.9                 |
| Σ S-type$^b$      | 80.8 b              |
| November           |                     |
| M I               | 16.9 aB             |
| M II              | 16.6 B              |
| M III             | 23.7 aB             |
| M IV              | 3.4 bA              |
| LSD$_{0.05}$      | 9.9                 |
| Σ M-type$^b$      | 9.3                 |
| Σ S-type$^b$      | 61.0 a              |

| 2015               |                     |
| September          |                     |
| M I               | 5.7 aA              |
| M II              | 17.1 B              |
| M III             | 22.9 bB             |
| M IV              | 5.7 aA              |
| LSD$_{0.05}$      | 10.8                 |
| Σ M-type$^b$      | 10.9                 |
| Σ S-type$^b$      | 51.4 a              |
| October            |                     |
| M I               | 29.4 bB             |
| M II              | 14.7 A              |
| M III             | 11.8 aA             |
| M IV              | 5.9 aA              |
| LSD$_{0.05}$      | 9.4                 |
| Σ M-type$^b$      | 11.3                 |
| Σ S-type$^b$      | 61.8 b              |
| November           |                     |
| M I               | 9.7 aA              |
| M II              | 16.1 AB             |
| M III             | 12.9 abAB           |
| M IV              | 19.4 bB             |
| LSD$_{0.05}$      | 10.8                 |
| Σ M-type$^b$      | 9.3                 |
| Σ S-type$^b$      | 58.1 ab             |

| Overall month     |                     |
| September         |                     |
| M I               | 17.2 aB             |
| M II              | 18.2 B              |
| M III             | 35.4 bC             |
| M IV              | 1.9 aA              |
| LSD$_{0.05}$      | 11.6                 |
| Σ M-type$^b$      | 12.1                 |
| Σ S-type$^b$      | 72.7                |
| October           |                     |
| M I               | 24.2 bBC            |
| M II              | 15.4 B              |
| M III             | 30.9 abC            |
| M IV              | 2.0 aA              |
| LSD$_{0.05}$      | 10.8                 |
| Σ M-type$^b$      | 11.6                 |
| Σ S-type$^b$      | 72.5                |
| November          |                     |
| M I               | 22.9 abB            |
| M II              | 14.7 AB             |
| M III             | 20.4 aB             |
| M IV              | 8.3 bA              |
| LSD$_{0.05}$      | 4.2                 |
| Σ M-type$^b$      | 10.8                 |
| Σ S-type$^b$      | 66.4                |
| Overall year      |                     |
| 2013              |                     |
| M I               | 21.0 abB            |
| M II              | 18.7 B              |
| M III             | 41.8 bC             |
| M IV              | 0.7 aA              |
| LSD$_{0.05}$      | 11.6                 |
| Σ M-type$^b$      | 12.3                 |
| Σ S-type$^b$      | 82.2 b              |
| 2014              |                     |
| M I               | 28.4 bC             |
| M II              | 13.7 B              |
| M III             | 29.1 bC             |
| M IV              | 1.1 aA              |
| LSD$_{0.05}$      | 13.1                 |
| Σ M-type$^b$      | 9.3                 |
| Σ S-type$^b$      | 72.3 ab             |
| 2015              |                     |
| M I               | 14.9 a              |
| M II              | 16.0                 |
| M III             | 15.8 a              |
| M IV              | 10.3 b              |
| LSD$_{0.05}$      | 15.1                 |
| Σ M-type$^b$      | ns                  |
| Σ S-type$^b$      | 57.1 a              |
| Overall year+month|                     |
| 2013              |                     |
| M I               | 21.5 B              |
| M II              | 16.1 AB             |
| M III             | 28.9 B              |
| M IV              | 4.1 A               |
| LSD$_{0.05}$      | 15.6                 |
| Σ M-type$^b$      | 70.5                |
| Σ S-type$^b$      | 5.6 A               |

a Phenotypic features of mycelial types I-IV and sclerotial types I-IV are described in Table 2
b Σ M-type$^b$ = summarized distribution for mycelial types I-IV; Σ S-type$^b$ = summarized distribution for sclerotial types I-IV

For each column, different small letters indicate significant differences at $P = 0.05$ among years, months. For each row, different capital letters indicate significant differences at $P = 0.05$ among morphological types M or S.
Previous studies showed that an increased concentration of phenolic compounds inhibited the growth of *B. cinerea* (Padgett and Morrison 1990; Pezet et al. 2003; Deytieux-Belleau et al. 2009; Köycü et al. 2014). In addition, the growth and latency of *B. cinerea* were inhibited more in the berries of dark-skinned cultivars compared to white-skinned ones with lower concentration of phenolic compounds. In our study, although berries of dark-skinned cv ‘Turán’ contained more polyphenols (Balga et al. 2014) than the white-skinned cv ‘Olaszrizling’, differences in mycelial growth rate and morphological types of *B. cinerea* did not occurred between isolates originated from white- and dark-skinned cultivars (non-significant cultivar effect in ANOVA). This result clearly indicated that the effect of dark-skinned cultivars on gray mold inhibition might occur in the plant tissues (such as in berries) but did not cause differences in the growth and morphological features of *B. cinerea* isolates in the laboratory originated either from dark- or white-skinned cultivars.

This study was the first confirming any effects of sampling months on mycelial growth and morphological types of *B. cinerea* during noble rot development (Tables 5 and 6) and their possible connection with noble rot incidence in the field. Magyar and Bene (2006) described the morphological features of *B. cinerea* in noble rotted berries under the specific weather conditions of the Tokaj wine district but these features were not connected further to sampling months. Our study showed that the effect of sampling months had stronger effects on the morphological types of *B. cinerea* isolates than on the mycelial growth rate of the isolates (Tables 5 and 6), which may influence the botrytisation process of berries in earlier and later months of the autumn.

In conclusion, our results showed possible effects of year/sampling month on noble rot incidence in connection with mycelial growth rate and morphological types of *B. cinerea* strains isolated during noble rot development. These differences may also appear in other European vineyards during the noble rot development of berries in autumn.

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**Compliance with ethical standards**

**Conflict of interest** The authors declare no conflict of interest.

**Ethical approval** This manuscript is original and not published elsewhere. The authors discussed the result, read and approved the final article. The authors confirm that there are no ethical issues in publication of the manuscript.

**Research involving human participants and/or animals** This study does not contain studies with human participants or animals performed by any of the authors.

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