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- Efficiency of bio-fungicides (Trichoderma spp and Pseudomonas fluorescens) on
- 4 seedling emergence, vigour and health of infected chilli seeds (*Capsicum annuum*)
- 5 by Colletotrichum capsici

6 ABSTRACT

Damping off and fruit rot caused by *Colletotrichum capsici* are the major constraints in production and marketability of chilli. Systemic fungicides are commonly used to control this disease. However, continuous use of chemical fungicides leads to negative impact on environment, soil and human health. Therefore, present studies (blotter and pot experiment) were conducted to explore the biofungicides (as an alternative to chemical fungicide) in comparison with carbendazim using chilli seeds infected with *Colletotrichum capsici*.

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Experiments were conducted at the CCSHAU, Hisar, India during 2016 in completely randomized 14 15 design with nine treatments replicated three times. Six months old seeds having germination above the Indian Minimum Seed Certification Standard, were infected with Colletotrichum capsici and such 16 infected seeds were treated with Trichoderma asperellum, Trichoderma viridae, Pseudomonas 17 fluorescens individually and their combinations to control the disease incidence. The infected, un-18 19 infected and seed treatment with carbendazim served as controls. Results revealed that the seed 20 germination was significantly high (94.7 %) with Trichoderma viride treatment as compared to all other treatments including controls in blotter method. However, the seedling emergence in pot culture was 21 22 significantly superior with Carbendazim treatment, the seed treatment with Pseudomonas fluorescens 23 and Trichoderma viridae was on par to that of Carbendazim treatment. The seedling length was significantly superior with Trichoderma viride compared to the carbendazim and other controls both in 24 25 blotter and pot culture. The seedling dry weight and seedling vigour were significantly high with 26 carbendazim as compared to the Trichoderma viride treatment or other treatments in both blotter and 27 pot culture. However, the overall seedling vigour obtained with Trichoderma viride was same to that of 28 carbendazim treatment. The disease incidence was significantly least with Pseudomonas fluorescens 29 as compared to the Trichoderma viride and carbendazim in blotter method and; T. viride + P. 30 fluorescens treatment was on par to that of carbendazim treatment in pot culture. Therefore, use of 31 Trichoderma viride and Pseudomonas fluorescens individually or in combinations are suggested as an alternative to carbendazim to control the Colletotrichum capsici. 32

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34 **Key words:** Chilli; *Colletotrichum capsici;* Carbendazim; bio-fungicides.

35 1. INTRODUCTION

Chilli is a major spice crop in India and India stands 3rd in production (21). The crop is suffered mainly 36 37 by seedling rot and fruit rot caused by Colletotrichum capsici leading to reduced marketability and fruit 38 yield [14, 15]. To control this disease, systemic fungicides are commonly used, especially the 39 carbendazim at the recommended dose of 0.2 % [17]. However, continuous use of chemical 40 fungicides has deleterious effects on biodiversity, environment and human health [3]. In this direction, 41 several reports show the effect of bio-fungicides like, Trichoderma viride, Pseudomonas fluorescens 42 etc. on control of Colletotrichum capsici and improved the seedling parameters and yield of chilli with 43 a decreased fruit rot [8, 11, 23, 24, 25, 26]. As most of the studies pertain only to bio-fungicides, it is 44 pertinent to identify a bio-fungicide comparable to that of chemical fungicides in the changing climate 45 scenario. Hence, the present study was undertaken to study the effect of bio-fungicides viz., 46 Trichoderma viride, Pseudomonas fluorescens, Trichoderma asperellum individually and their 47 combinations in comparison with chemical fungicide (carbendazim), infected and un-infected seed on

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48 seed germination, seedling emergence, seedling vigour and disease control in chilli seeds infected 49 with Colletotrichum capsici.

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51 2. MATERIAL AND METHODS

52 Two experiments (blotter and pot culture) were conducted to study the effect of bio-fungicides on 53 seed quality parameters of chilli seeds infected with Colletotrichum capsici. These experiments were 54 conducted at the Department of Seed Science and Technology, CCSHAU, Hisar during October-55 November, 2016. The seeds used in these experiments were six months old which were harvested 56 during February - March, 2016 (high yielding popular variety, RCH-1). The seed germination was above the Indian Minimum Seed Certification Standards. 57

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The experiments were conducted with nine treatments in three replications (Table 1) both in blotter 59 and pot experiments. In blotter method, petri dishes(15 cm diameter) lined with two layers of blotting 60 paper (Whatman No.1) were prepared, adequately watered, 25 seeds in each petri dish were placed 61 and kept in BOD (biological oxygen demand) incubator for 14 days at 25°C. Sixteen petri plates were 62 used for each replication. These plates were watered as and when the blotter paper appeared nearly 63 64 to dryness. For pot culture experiment, pots (27.5 cm diameter and 30 cm height) were filled with four 65 kg of oven sterilized soil. The soil is sandy loam with organic carbon (0.15%), pH (8.1) and Ec (0.15 66 dS/m at 25^oC) [22]. Twenty five seeds were placed at a depth of 1-2 cm in each pot and eight pots 67 replication were maintained. The pots were watered daily up to 14 days. The weeds were uprooted 68 whenever appeared.

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70 In both the experiments, final germination count, disease incidence and disease control was monitored on 14th day, and ten randomly selected seedlings per replication were taken for observations on shoot 71 length, root length and total seedling length. After taking the shoot and root length, the same seedlings 72 were kept for drying in oven at 70±1°C until they attained a constant dry weight and calculated the 73 74 seedling vigour. The formulae used for various calculations are given below [1].

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76 Table 1. Treatment details involved in the experiment

No.	Treatments
T ₁	Trichoderma viride (200 mg/ 20 g seed)
T_2	Trichoderma asperellum (200 mg/ 20 g seed)
T ₃	Pseudomonas fluorescens (200 mg/ 20 g seed)
T ₄	Trichoderma asperellum (100 mg/ 20 g seed)+ Trichoderma viride (100 mg/ 20 g seed)
T ₅	Pseudomonas fluorescens (100 mg/ 20 g seed) + Trichoderma viride (100 mg/ 20 g seed)
Т ₆	Pseudomonas fluorescens (100 mg/ 20 g seed) + Trichoderma asperellum (100 mg/ 20 g seed)
T ₇	Infected seed (<i>Colletotrichum capsici</i> infected seed but not treated with bio-fungicide or Carbendazim)
T ₈	Un-infected seed (Six months old seed which was not-infected with <i>Colletotrichum capsici</i> and not treated with bio-fungicide or Carbendazim)
Т ₉	Carbendazim treated (<i>Colletotrichum capsici</i> infected seed treated with Carbendazim (40 mg/ 20 g seed)
Note:	(1) Dose:10 g kg ⁻¹ alone and in combination @ 5 + 5 g kg ⁻¹ seed, (2) Except the controls (T_7

77 78 and T₈), all the bio-fungicides and Carbendazim treatments were given to the seeds that are infected

79 with Colletotrichum capsici.

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81 Isolation of Colletotrichum capsici and seed infection

The infected chilli fruit portion was sterilized and cultured on potato dextrose agar (PDA) medium in a petri plate. The pure culture of *Colletotrichum capsici* was identified, isolated, sub-cultured and multiplied again on PDA and used for seed infection. The multiplication has taken 7-9 days.

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Twenty gram of chilli seeds were taken in a beaker. The *Colletotrichum capsici* was scrapped into the beaker containing the seeds using a scrapper in the laminar air flow. The beaker was closed with para-film tape, shaken for 15 minutes and left undisturbed for 24h. Such infected seeds with *Colletotrichum capsici* were used further for treatment with bio-fungicides or carbendazim.

90 91 Seed treatment

Seed treatment with bio-fungicide or carbendazim 92 The Colletotrichum capsici infected seeds were treated with different bio-fungicides, Trichoderma 93 asperellum, Trichoderma viridae and Pseudomonas fluorescens (200 mg/ 20 g seed) individually and 94 in combinations (100mg + 100mg) or with carbendazim (40 mg/ 20 g seed) in a beaker, shaken gently 95 to cover the seed uniformly with bio-fungicide or carbendazim (Table 1). The control treatments were, 96 infected seed (not treated with any bio-fungicide or Carbendazim), uninfected seed (six months old 97 seed which was not treated with any bio-fungicide or Carbendazim) and Carbendazim treatment 98 (infected seed treated with Carbendazim). 99

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100	Number of seeds germinated
101	Seed germination (%) in blotter = × 100
102 103	Total number of seeds placed for germination
103	Number of seedlings emerged
105	Seedling emergence (%) in pot = × 100
106	Total number of seeds placed for seedling emergence
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109	Seedling length (cm) = Seedling shoot length (cm) + Seedling root length (cm)
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111	Seed Vigour Index I = Seed germination percentage × Seedling length (cm) in blotter method
112 113	Seed Vigour Index I = Seedling emergence percentage × Seedling length (cm) in pot culture
115	Seed vigour index i = Seeding emergence percentage x Seeding length (cm) in pot culture
115	Seed vigour Index II = Seed germination percentage × Dry seedling weight (mg) in blotter method
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117	Seed vigour Index II = Seed emergence percentage × Dry seedling weight (mg) in pot culture
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119	
120	Number of seedlings affected either in blotter or pot
121	Disease incidence (%) =
122	Total number of seedlings either in blotter or pot
123 124	Treatment - Infected
124	Disease control (%) = × 100
125	Treatment
127	
128	Where, treatment refers to all the eight treatments including two controls namely, un-infected
129	and Carbendazim treatments.
120	The data obtained was statistically analyzed in Completely Rendemized Design (CRD) in both

130 The data obtained was statistically analyzed in Completely Randomized Design (CRD) in both 131 the experiments.

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133 3. RESULTS AND DISCUSSION

134 **3.1 Seed germination**

The seed germination was significantly superior in blotter method (87.6 %) as compared to the seedling emergence in the pot experiment (84.5 %) although the differences are marginal (3.5 %). In blotter method among the treatments only *T. viride* (94.7 %) showed significantly higher seed germination compared to all other treatments including the carbendazim treatment (92.0 %). While, *Pseudomonas fluorescens* treatment (92.7 %) was on par to the carbendazim treatment (Table 2). Although the differences between the treatments are meagre, the germination percentage was markedly high both in the bio-fungicide treatments and carbendazim treatment compared to the controls (infected seed and un-infected seed). The higher seed germination with bio-fungicides could be through inhibition of growth of *C. capsici* [18, 27].

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145Table 2. Effect of bio-fungicides on seed germination, seedling length and dry weight in chilli146seeds infected with Colletotrichum capsici

Treatments	Seed germinati on (%)	Seedling emergen ce (%)	Seedling length (cm		Seedling dry weight (mg/ seedling)	
	Blotter	Pot	Blotter	Pot	Blotter	Pot
Trichoderma viride	94.7 (76.6) ^g	89.0 (70.6) ^a	7.04 ^d	7.45 ^d	32.21 ^d	30.00 ^d
Trichoderma asperellum	86.7 (68.6) ^f	84.3 (66.7) ^b	4.96 ^a	6.76 ^a	29.23 ^b	32.23 ^d
Pseudomonas fluorescens	92.7 (74.3) ^a	83.3 (65.9) ^b	4.89 ^a	7.01 ^a	28.50 ^b	30.16 ^d
Trichoderma asperellum + Trichoderma viride	88.0 (69.7) ^e	85.3 (67.5) ^b	5.66 ^b	6.60 ^a	27.83 ^b	29.86 ^d
Pseudomonas fluorescens + Trichoderma viride	89.3 (70.9) ^d	89.3 (71.0) ^a	4.90 ^a	6.64 ^a	33.56 ^d	30.20 ^d
Pseudomonas fluorescens + Trichoderma asperellum	89.3 (70.9) ^d	85.0 (67.2) ^b	4.87 ^c	7.09 ^a	32.67 ^d	35.76 ^b
Infected seed	70.3 (57.0) ^c	69.3 (56.4) ^c	4.69 ^c	4.58 ^c	25.67 ^c	25.66 ^c
Un-infected seed	85.3 (67.5) ^b	83.7 (66.1) ^b	5.94 ^b	5.22 ^b	28.00 ^b	34.33 ^b
Carbendazim treated seed	92.0 (73.5) ^a	91.0 (72.5) ^a	5.26 ^ª	6.68 ^ª	36.67ª	39.33ª
Mean	87.6 (69.9)	84.5 (67.1)	5.36	6.45	30.48	31.95
C.D (P< 0.05)	0.8	1.8	0.38	0.43	1.36	2.40
SEm <u>+</u>	0.3	0.6	0.13	0.14	0.45	0.80
C.V. (%)	0.7	1.6	4.19	3.87	2.58	4.35

¹⁴⁷ Note: Values in parenthesis are arc sign transformed values for statistical analyses

148 In pot culture experiment, the seedling emergence was significantly superior with carbendazim as 149 compared to all the bio-fungicide treatments and other control treatments. However, the seed 150 germination was above the minimum standards of seed germination in all the treatments except the infected seeds (absolute control) both in blotter and pot culture experiments. Hence, for the purpose of 151 higher seed germination any of the bio-fungicides may be suggested to achieve higher seed 152 153 germination or seedling emergence of chilli seeds. Both in blotter and pot culture, infected seed 154 maintained showed significantly lower seed germination and seedling emergence respectively as 155 compared to the un-infected control or carbendazim treatments (Table 2).

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157 **3.2 Seed quality parameters**

Both in blotter and pot culture, among the treatments, seedling length was significantly superior with *Trichoderma viride* (7.04 cm in blotter and 7.75 cm in pot culture) as compared to the carbendazim 160 (5.26 and 6.68 cm respectively). In pot culture, bio-fungicide treatments showed significantly higher 161 seedling length as compared to the un-infected seed. Similar results of increased seedling length due 162 to application of *Trichoderma viride*, *Trichoderma asperellum* and *Pseudomonas fluorescens* 163 individually or in combination was reported in different species [7,19, 20].

164 Seedling dry weight among the treatments and across the two experiments was significantly higher in 165 carbendazim treated seeds (36.67 mg in blotter method and 39.33 mg in pot culture) as compared to all bio-fungicide treatments (Table 2). However, in pot culture, bio-fungicide treatments performed 166 167 better over the un-infected seed, this could be due to effective control of pre-emergence and post-168 emergence damping off through decreased colony formation by C. capsici [10, 12]. In pot culture 169 (similar to field conditions), lower effect of bio-fungicides could be due to longer time required for 170 perpetuation of bio-fungicides in view of requirement of carbohydrate at early stages, whereas, 171 carbendazim do not depend on seedling for carbohydrate requirement.

172 3.3 Seedling vigour

173 Seedling vigour is an important trait in ensuring proper crop establishment and economic yields especially under adverse conditions. Seed borne pathogen like C. capsici is known to affect the 174 seedling vigour causing fruit rot and reduces the yield. Under such conditions, application of chemical 175 176 fungicide or bio-fungicide would help to combat the effects of C. capsici. Several reports have shown 177 the positive influence of bio-fungicides like Trichoderma and others on seedling vigour in chilli [2, 9, 178 18, and 19]. However, scanty literature is available with respect to comparison of bio-fungicides with 179 carbendazim which is a popular systemic fungicide [17]. Therefore, it is very pertinent to identify a bio-180 fungicide comparable to that of carbendazim in the changing climate scenario as carbendazim have 181 deleterious effects on biodiversity, environment and human [3]. In the present study, seedling vigour 182 index-I and II were significantly high with carbendazim treatment compared to all the bio-fundicides 183 and control treatments in both blotter and pot culture (Table 3). Further, all the bio-fungicide 184 treatments found superior over the un-infected seed for SVI-I in pot culture and SVI-II in blotter 185 method (Table 3). These differences are due to variations in seed germination, seedling length and 186 seedling dry weights in calculation of seedling vigour indices. However, when the data was 187 normalized by giving equal weightage to unity for all three parameters, seedling vigour with 188 Trichoderma viride found on par to the carbendazim treatment (Table 3). Similarly, Choudhary et al. 189 [5] reported that Trichoderma viride was effective as compared to the carbendazim in terms of 190 seedling vigour. Further, all the bio-fungicides were better than the control (un-infected seed). Hence, 191 seed treatment with Trichoderma viride is suggested to combat the C. capsici and thus to achieve 192 healthy vigorous seedlings for better yields of chilli.

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Table 3. Effect of bio-fungicides on seedling vigour index in *chilli* seeds infected with *Colletotrichum capsici*

Treatments	SVI-I		SVI-II		Overall SVI
	Blotter	Pot	Blotter	Pot	Pooled
Trichoderma viride	498.6 ^b	601.6 ^{ad}	3048.8 ^g	2670.0 ^{def}	0.81
Trichoderma asperellum	430.1 ^d	628.9 ^ª	2533.6 ^{ef}	2718.4 ^{befg}	0.60
Pseudomonas fluorescens	453.8 ^d	584.2 ^{ad}	2641.2 ^e	2513.8 ^d	0.59
Trichoderma asperellum + Trichoderma viride	498.1 ^b	563.5 ^d	2449.3 ^{bf}	2544.4 ^{dg}	0.60
Pseudomonas fluorescens + Trichoderma viride	438.4 ^d	593.4 ^{ad}	2997.9 ^{dg}	2698.9 ^{bf}	0.64
Pseudomonas fluorescens + Trichoderma asperellum	435.7 ^d	603.0 ^{ad}	2918.3 ^d	3040.0 ^b	0.69
Infected seed	329.8 [°]	317.5 [°]	1805.3 [℃]	1779.3 ^c	0.33
Un-infected seed	506.9 ^b	436.7 ^b	2389.0 ^b	2873.0 ^b	0.57
Carbendazim treated seed	647.7 ^a	608.5 ^ª	3373.3ª	3579.3ª	0.81

Mean	471.0	548.6	2684.1	2713.0
C.D (P< 0.05)	32.83	40.05	127.27	198.24
SEm <u>+</u>	10.96	13.37	42.50	66.21
C.V. (%)	4.03	4.22	2.74	4.22

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197 **3.4 Disease infection and disease control**

198 In blotter experiment, disease incidence was significantly less in Pseudomonas fluorescens (5.33 %) as compared to the carbendazim (8.00 %), whereas, the Trichoderma viride (7.33 %) was comparable 199 200 to the carbendazim (Table 4). In pot culture, carbendazim showed significantly lower disease 201 incidence (9.0 %) but was on par to that of Trichoderma viride (11.0%) and Trichoderma viride + Pseudomonas fluorescens (10.67 %). All bio-fungicide treatments resulted in significantly lower 202 203 disease incidence or on par to the un-infected seed (control) (Table 4). In contrast to disease incidence, the disease control was significantly higher in Pseudomonas fluorescens (81.61 %) as 204 205 compared to carbendazim (73.03 %) and Trichoderma viride (74.71 %) in blotter technique (Table 4). 206 In pot culture, disease control was significantly superior in carbendazim (70.65 %) treatment compared to all bio-fungicides except Trichoderma viride + Pseudomonas fluorescens (64.44 %). 207

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Table 4. Effect of bio-fungicides on disease incidence and disease control in *chilli* seeds infected with *Colletotrichum capsici*

Treatments	Disease ind	cidence (%)	Disease control (%)		
	Blotter	Pot	Blotter	Pot	
Trichoderma viride	7.33 (15.70) ^a	11.00 (19.36) ^d	74.71 (59.80) ^a	63.33 (52.71) ^d	
Trichoderma asperellum	13.33 (21.41) ^b	15.67 (23.31) ^b	54.02 (47.29) ^g	47.78 (43.71) ^b	
Pseudomonas fluorescens	5.33 (13.34) ^b	16.67 (24.08) ^b	81.61 (64.60) ^f	44.44 (41.79) ^b	
Trichoderma asperellum + Trichoderma viride	12.00 (20.26) ^e	14.67 (22.47) ^b	58.62 (49.94) ^e	51.11 (45.63) ^b	
Pseudomonas fluorescens + Trichoderma viride	10.67 (19.05) ^d	10.67 (18.98) ^a	63.22 (52.65) ^d	64.44 (53.46) ^{ad}	
Pseudomonas fluorescens + Trichoderma asperellum	10.67 (19.05) ^d	15.00 (22.77) ^b	63.22 (52.65) ^d	50.00 (44.98) ^b	
Infected seed	29.67 (32.99) ^c	30.67 (33.61) ^c	0.00 (0.00) ^c	0.00 (0.00) ^c	
Un-infected seed	14.67 (22.51) ^b	16.33 (23.83) ^b	50.57 (45.31) ^b	46.74 (43.11) ^b	
Carbendazim treated seed	8.00 (16.42) ^a	9.00 (17.45) ^a	73.03 (58.69) ^a	70.65 (57.17) ^a	
Mean	12.41 (15.52)	20.08 (22.87)	57.67 (47.88)	48.72 (42.51)	
C.D (P< 0.05)	0.83	1.82	1.67	4.03	
SEm <u>+</u>	0.28	0.61	0.56	1.34	
C.V. (%)	2.40	4.59	2.02	5.48	

211 Note: Values in parenthesis are arc sign transformed values for statistical analyses

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Many reports have shown that the bio-fungicides like *Trichoderma viride*, *Pseudomonas fluorescens* and their combinations inhibited the mycelia growth of pathogen and hence disease control caused by *C. capsici* [8, 11, 23, 24, 25, 26]. These studies have not compared the effectiveness of bio-fungicide against the carbendazim which is a popular systemic fungicide. However, a few studies show that chemical fungicides like copper oxychloride is more effective than *Trichoderma viride* in controlling the disease caused by *C. capsici* [16]. The bio-fungicide, *Trichoderma viride* produce antibiotic (trichodermin) and extracellular enzymes (chitinase, cellulose) those inhibit the plant pathogen [19]. Further, it was effective with combined use of bio-fungicide and carbendazim in reducing the disease incidence, thus higher yield and quality of chilli was achieved [6, 13]. Further, both seed treatment and soil treatment are suggested for effective control of *C. capsici* [4].

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Therefore, the use of *Trichoderma viride* and *Pseudomonas fluorescens* or their combinations are suggested in place of carbendazim against *Colletotrichum capsici* and for better seed quality parameters in chilli.

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229 4. CONCLUSION

Seed treatment with *Trichoderma viride* (10g kg⁻¹ seed) and *Pseudomonas fluorescens* (10g kg⁻¹ seed) individually or combination (*Trichoderma viride, 5*g kg⁻¹ seed + *Pseudomonas fluorescens, 5*g kg⁻¹ seed) can be effectively used in place of carbendazim (0.2 %) treatment for effective control of *Colletotrichum capsici* to achieve higher seedling vigour.

235 COMPETING INTERESTS

237 Authors declare no competing interest exists.

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