

1 Bimatrix games that include interaction times alter the
2 evolutionary outcome: The Owner–Intruder game

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9 **Abstract**

Classic bimatrix games, that are based on pair-wise interactions between two opponents in two different roles, do not consider the effect that interaction duration has on payoffs. However, interactions between different strategies often take different amounts of time. In this article, we further develop a new approach to an old idea that opportunity costs lost while engaged in an interaction affect individual fitness. We consider two scenarios: (i) individuals pair instantaneously so that there are no searchers, and (ii) searching for a partner takes positive time and populations consist of a mixture of singles and pairs. We describe pair dynamics and calculate fitnesses of each strategy for a two-strategy bimatrix game that includes interaction times. Assuming that distribution of pairs (and singles) evolves on a faster time scale than evolutionary dynamics described by the replicator equation, we analyze the Nash equilibria (NE) of the time-constrained game. This general approach is then applied to the Owner–Intruder bimatrix game where the two strategies are Hawk and Dove in both roles. While the classic Owner–Intruder game has at most one interior NE and it is unstable with respect to replicator dynamics, differences in pair duration change this prediction in that up to four interior NE may exist with their stability depending on whether pairing is instantaneous or not. The classic game has either one (all Hawk) or two ((Hawk,Dove) and (Dove,Hawk)) stable boundary NE. When interaction times are included, other combinations of stable boundary NE are possible. For example, (Dove,Dove), (Dove,Hawk), or (Hawk,Dove) can be the unique (stable) NE if interaction time between two Doves is short compared to some other interactions involving Doves.

10 *Keywords:* Evolutionary game theory, Hawk–Dove game, Pair formation, Time-scale
11 separation, Time-constrained games

12 1. Introduction

13 Classic evolutionary game theoretical models in normal form consider two players with
14 a finite number of strategies and a payoff matrix. Players in a large (infinite) population
15 meet at random, interact pair-wise, and obtain their corresponding (individual) fitnesses.
16 There are three important and somewhat hidden assumptions: (i) interaction times be-
17 tween two strategies are not considered, i.e., they are all assumed to be the same, (ii) the
18 distribution of strategy pairs corresponds to random pair formation among all individuals
19 and (iii) individual fitness accrues only through pair interactions. These assumptions fit
20 genetic population models with two (or more) alleles at a single locus. In the genetic
21 model, the alleles pair randomly during meiosis and the resulting distribution of geno-
22 types is given by the Hardy–Weinberg equation. When alone, alleles cannot gain any
23 fitness. For many phenotypic models (e.g., the Hawk–Dove, or Prisoner’s dilemma), these
24 assumptions are likely not satisfied. For example, when two aggressive individuals are in
25 a fight, their interaction can be much longer when compared to the situation where one
26 individual (a Dove) exits from an interaction with a Hawk (in which case the Hawk will
27 win the contest). Because contests between different strategies can take different times,
28 the resulting equilibrium distribution of pairs does not correspond to the Hardy–Weinberg
29 equation.

30 Křivan and Cressman (2017) showed that, when individuals pair instantaneously but
31 the interaction times are strategy dependent, the Hawk–Dove model may have a mixed
32 ESS (i.e., an evolutionarily stable state that consists of a mixture of Hawks and Doves)
33 when the cost of a fight is lower than the value of the contested resource. For this
34 to happen, the interaction time between two Hawks must be long enough relative to
35 interaction times between other strategies. Such an outcome is not possible in the classic
36 Hawk–Dove game that does not consider interaction times. Similarly, for the repeated
37 Prisoner’s dilemma, provided cooperators stay together for enough rounds of the game
38 while pairs with at least one defector disband quickly, cooperation does evolve (Křivan
39 and Cressman, 2017). This situation arises naturally if players can choose whether to
40 continue the game to the next round with the same opponent, since it is always better to
41 play against a cooperator than a defector in the Prisoner’s dilemma game (see also the
42 opting-out game (Zhang et al., 2016)).

43 Moreover, individuals can gain/lose fitness when alone (e.g., individuals with different
44 strategies may have different mortalities). While the above games do not consider singles,
45 Křivan et al. (2018) assumed that pairing between individuals is not immediate and being
46 single has fitness consequences. They showed that distributional dynamics alone can lead
47 to density dependence in models (e.g., the Hawk–Dove model) that are only frequency
48 dependent when pairing is instantaneous and all interaction times are the same.

49 All the models considered above are based on symmetric games (in particular, matrix
50 games), where the two contestants are assumed to be drawn from the same population
51 and can differ only in their choice of strategy. It is well known that various asymmetries
52 (Broom and Rychtář, 2013) in contestants lead to qualitatively different outcomes when
53 interaction times are not considered. A class of asymmetric games, bimatrix games,
54 where the two contestants are drawn from two different types of individuals (e.g., two
55 populations or two roles) was studied thoroughly in the literature (e.g., Hofbauer and
56 Sigmund, 1998; Cressman, 2003; Broom and Rychtář, 2013). A well-known result of
57 classic evolutionary game theory for these games is that no interior evolutionarily stable
58 strategy exists (Selten, 1980) (i.e., no ESS where each population is a mixture of pure
59 strategies). Furthermore, bimatrix games may have an interior Nash equilibrium (NE) but
60 it cannot be asymptotically stable under the (bimatrix) replicator equation, the standard
61 game dynamics of evolutionary game theory (Hofbauer and Sigmund, 1998). In particular,
62 ESSs and asymptotically stable equilibria correspond to strict NEs of the bimatrix game
63 (i.e., pure strategy pairs where both players do strictly worse by unilaterally changing
64 their strategy).

65 Given the conceptual differences between the evolutionary outcomes of classic matrix
66 and bimatrix games, it is important to understand the consequences of strategy-dependent
67 interaction times by extending the analysis beyond the matrix games considered by Krivan
68 and Cressman (2017). To this end, in this article, we study the effect of interaction time on
69 the evolutionary outcome of bimatrix games when both populations have two strategies.
70 We consider two pair formation processes based on the assumption that the number of
71 individuals of each population are the same. In Section 2, as existing pairs disband,
72 these individuals instantaneously form new pairs randomly among themselves. From the
73 analytic expression of the equilibrium distribution of pairs at a given number of each
74 strategy in both populations, we analyze the resulting game (i.e., investigate its NEs
75 and their stability) when individual fitness is defined as expected payoff per unit time.
76 When interaction times are all the same, we recover the classic results. Otherwise, more
77 complicated evolutionary outcomes emerge such as multiple interior NEs (some of which
78 are stable and some unstable) as well as strict NE that differ from the classic game.
79 These possibilities are illustrated there by a thorough analysis of the Owner–Intruder
80 game (Broom and Rychtář, 2013), the bimatrix version of the Hawk–Dove game where
81 individuals assume one of the two roles, owner or intruder.

82 In Section 3, when pairs disband, the resulting singles form new pairs at random
83 through the mass action principle with a finite encounter rate. Since the analytic expres-
84 sion of the equilibrium distribution of pairs at a given number of each strategy in both
85 populations is no longer tractable unless all interaction times are the same, we analyze
86 the Owner–Intruder game, with unequal interaction times, numerically.

87 **2. Instantaneous pair formation**

88 We consider a bimatrix game with two strategies denoted by e_i ($i = 1, 2$) for the row
 89 player in population 1 and f_j ($j = 1, 2$) for the column player in population 2. The payoff
 90 bimatrix is

$$\begin{array}{cc} & f_1 & f_2 \\ e_1 & \left[\pi_{11}^e, \pi_{11}^f & \pi_{12}^e, \pi_{12}^f \right] \\ e_2 & \left[\pi_{21}^e, \pi_{21}^f & \pi_{22}^e, \pi_{22}^f \right] \end{array} \quad (1)$$

91 where π_{ij}^e (respectively, π_{ij}^f) is the payoff to e_i (respectively f_j) when interacting with f_j
 92 (respectively e_i). In contrast to classic evolutionary game theory, we explicitly incorporate
 93 the duration of interactions into the game through the time interaction matrix

$$\begin{array}{cc} & f_1 & f_2 \\ e_1 & \left(\tau_{11} & \tau_{12} \right) \\ e_2 & \left(\tau_{21} & \tau_{22} \right) \end{array} \quad (2)$$

94 where τ_{ij} is the expected time two players using strategy e_i and f_j stay together.

95 In this section, we assume that, when pairs split, all these newly single individuals
 96 immediately form new pairs at random. We are interested in the equilibrium distribution
 97 of strategy pairs (e_i, f_j) for given numbers of the different strategies. Let n_{ij} be the
 98 number of strategy pair (e_i, f_j) . As shown in Appendix A, pair dynamics are

$$\begin{aligned} \frac{dn_{11}}{dt} &= -\frac{n_{11}}{\tau_{11}} + \frac{\left(\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}}\right) \left(\frac{n_{11}}{\tau_{11}} + \frac{n_{21}}{\tau_{21}}\right)}{\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}} + \frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}}} \\ \frac{dn_{12}}{dt} &= -\frac{n_{12}}{\tau_{12}} + \frac{\left(\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}}\right) \left(\frac{n_{12}}{\tau_{12}} + \frac{n_{22}}{\tau_{22}}\right)}{\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}} + \frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}}} \\ \frac{dn_{21}}{dt} &= -\frac{n_{21}}{\tau_{21}} + \frac{\left(\frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}}\right) \left(\frac{n_{11}}{\tau_{11}} + \frac{n_{21}}{\tau_{21}}\right)}{\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}} + \frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}}} \\ \frac{dn_{22}}{dt} &= -\frac{n_{22}}{\tau_{22}} + \frac{\left(\frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}}\right) \left(\frac{n_{12}}{\tau_{12}} + \frac{n_{22}}{\tau_{22}}\right)}{\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}} + \frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}}} \end{aligned} \quad (3)$$

99 and the equilibrium distribution satisfies

$$\frac{n_{ij}}{\tau_{ij}} = \frac{\left(\frac{n_{i1}}{\tau_{i1}} + \frac{n_{i2}}{\tau_{i2}}\right) \left(\frac{n_{1j}}{\tau_{1j}} + \frac{n_{2j}}{\tau_{2j}}\right)}{\left(\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}} + \frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}}\right)} \quad \text{for } i, j = 1, 2. \quad (4)$$

100 Intuitively, at equilibrium, the number of disbanding (e_i, f_j) pairs per unit time (i.e., the
 101 left-hand side $\frac{n_{ij}}{\tau_{ij}}$ of (4)) must equal the number of newly formed (e_i, f_j) pairs from the
 102 newly single e_i strategists $\left(\frac{n_{i1}}{\tau_{i1}} + \frac{n_{i2}}{\tau_{i2}}\right)$ and f_j strategists $\left(\frac{n_{1j}}{\tau_{1j}} + \frac{n_{2j}}{\tau_{2j}}\right)$.

103 We observe that at the equilibrium distribution, $\frac{n_{ij}}{\tau_{ij}}$ satisfy the generalized Hardy–
 104 Weinberg equation, i. e.,

$$\frac{n_{11}}{\tau_{11}} \frac{n_{22}}{\tau_{22}} = \frac{n_{12}}{\tau_{12}} \frac{n_{21}}{\tau_{21}}. \quad (5)$$

105 Given the number of e_1 and f_1 strategists ($N_{e_1} = n_{11} + n_{12}$ and $N_{f_1} = n_{11} + n_{21}$, respec-
 106 tively) as well as the total number of individuals $N = n_{11} + n_{12} + n_{21} + n_{22}$ in either
 107 population, Appendix A shows that the unique nonnegative solution to (4) and (5) is
 108 (assuming $\tau_{12}\tau_{21} \neq \tau_{11}\tau_{22}$)

$$\begin{aligned} n_{11} &= \frac{\sqrt{A} + (N_{e_1} + N_{f_1})(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}) - N\tau_{12}\tau_{21}}{2(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})}, \\ n_{12} &= \frac{-\sqrt{A} + (N_{e_1} - N_{f_1})(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}) + N\tau_{12}\tau_{21}}{2(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})}, \\ n_{21} &= \frac{-\sqrt{A} - (N_{e_1} - N_{f_1})(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}) + N\tau_{12}\tau_{21}}{2(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})}, \\ n_{22} &= \frac{\sqrt{A} - (N_{e_1} + N_{f_1})(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}) + N(\tau_{12}\tau_{21} - 2\tau_{11}\tau_{22})}{2(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})}, \end{aligned} \quad (6)$$

109 where

$$A = (N\tau_{12}\tau_{21} - (N_{e_1} + N_{f_1})(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}))^2 + 4N_{e_1}N_{f_1}\tau_{11}\tau_{22}(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}). \quad (7)$$

110 When $\tau_{12}\tau_{21} = \tau_{11}\tau_{22}$ the above distributional equilibrium corresponds to the standard
 111 Hardy–Weinberg distribution

$$(n_{11}, n_{12}, n_{21}, n_{22}) = \left(\frac{N_{e_1}N_{f_1}}{N}, \frac{N_{e_1}N_{f_2}}{N}, \frac{N_{e_2}N_{f_1}}{N}, \frac{N_{e_2}N_{f_2}}{N} \right) \quad (8)$$

112 where $N_{e_2} \equiv N - N_{e_1}$ and $N_{f_2} \equiv N - N_{f_1}$. This is an important special case since it
 113 includes the classic situation, i.e., all interaction times are the same ($\tau_{11} = \tau_{12} = \tau_{21} = \tau_{22}$).

114 2.1. Fitness and evolutionary outcomes

115 Following Krivan and Cressman (2017), we define fitness as the expected payoff that
 116 an individual of a given phenotype obtains per unit of interaction time. For example, let

117 us consider an individual playing strategy e_1 in population 1. The probability that this
 118 individual is paired with an individual playing strategy f_1 is $n_{11}/(n_{11} + n_{12})$ and with an
 119 individual playing strategy f_2 is $n_{12}/(n_{11} + n_{12})$. When paired with an individual playing
 120 strategy f_1 , the focal individual receives payoff π_{11}^e/τ_{11} per unit of time. Similarly, when
 121 paired with an individual playing strategy f_2 , the focal individual gets payoff π_{12}^e/τ_{12} per
 122 unit of time. Thus, the focal individual has expected payoff (i.e., fitness) Π_{e_1} given by
 123 the first equation in (9). The fitness for individuals playing e_2 and those in the second
 124 population are calculated analogously, which leads to $(i, j = 1, 2)$

$$\begin{aligned}\Pi_{e_i} &= \frac{n_{i1}}{n_{i1} + n_{i2}} \frac{\pi_{i1}^e}{\tau_{i1}} + \frac{n_{i2}}{n_{i1} + n_{i2}} \frac{\pi_{i2}^e}{\tau_{i2}} \\ \Pi_{f_j} &= \frac{n_{1j}}{n_{1j} + n_{2j}} \frac{\pi_{1j}^f}{\tau_{1j}} + \frac{n_{2j}}{n_{1j} + n_{2j}} \frac{\pi_{2j}^f}{\tau_{2j}}.\end{aligned}\tag{9}$$

125 The corresponding time-constrained bimatrix game based on payoff bimatrix (1) and time
 126 interaction matrix (2) is then the two-strategy game with payoffs given by the fitness
 127 functions (9) evaluated at the distributional equilibrium (6) for fixed size N of each
 128 population.¹

129 To analyze this time-constrained bimatrix game, we examine how its NE structure
 130 depends on model parameters. We start by looking for NE in pure strategies (i.e., both
 131 populations are monomorphic) before considering NE where both populations are poly-
 132 morphic (i.e., the interior NE later in this section) and boundary NE (where exactly one
 133 population is polymorphic) in Section 2.3. Let us consider the equilibrium where all indi-
 134 viduals of population 1 play strategy e_1 while all individuals of the second population play
 135 strategy f_1 . Then $n_{11} = N$ and fitnesses of residents are $\Pi_{e_1} = \frac{\pi_{11}^e}{\tau_{11}}$ and $\Pi_{f_1} = \frac{\pi_{11}^f}{\tau_{11}}$. Now
 136 consider a mutant of the first population playing strategy e_2 in the resident system. This
 137 mutant can pair only with f_1 -strategists in which case its fitness is $\Pi_{e_2} = \frac{\pi_{21}^e}{\tau_{21}}$. Similarly,
 138 $\Pi_{f_2} = \frac{\pi_{12}^f}{\tau_{12}}$. Thus, the strategy (e_1, f_1) cannot be invaded if $\frac{\pi_{21}^e}{\tau_{21}} < \frac{\pi_{11}^e}{\tau_{11}}$ and $\frac{\pi_{12}^f}{\tau_{12}} < \frac{\pi_{11}^f}{\tau_{11}}$, in
 139 which case (e_1, f_1) is a strict NE.² Similar considerations for other pure strategy pairs
 140 show that a strategy (e_i, f_j) is a strict NE for the fitness functions given in (9) if it is a

¹We will use the phrase ‘‘fitness functions’’ rather than ‘‘payoffs’’ for these time-constrained games from now on to avoid confusion with payoffs in (1).

²If (e_1, f_1) is a strict NE, it must also resist invasion by mutants in population 1 that use any other strategy (including a mixed strategy) besides e_1 . However, since the fitness of the focal mutant is linear in the components of its mixed strategy, it is enough to verify (e_1, f_1) cannot be invaded by the pure strategy e_2 (and by f_2 in population 2).

141 strict NE of the classic game given by a time-adjusted payoff bimatrix

$$\begin{array}{c}
e_1 \\
e_2
\end{array}
\begin{array}{cc}
f_1 & f_2 \\
\left[\begin{array}{cc}
\frac{\pi_{11}^e}{\tau_{11}}, \frac{\pi_{11}^f}{\tau_{11}} & \frac{\pi_{12}^e}{\tau_{12}}, \frac{\pi_{12}^f}{\tau_{12}} \\
\frac{\pi_{21}^e}{\tau_{21}}, \frac{\pi_{21}^f}{\tau_{21}} & \frac{\pi_{22}^e}{\tau_{22}}, \frac{\pi_{22}^f}{\tau_{22}}
\end{array} \right]
\end{array}
. \quad (10)$$

142 We remark that the inequality conditions for a strict NE are independent of population
143 size. Furthermore, the fitness functions (9) when the populations are not monomorphic
144 are convex combinations of the appropriate entries in the time-adjusted payoff bimatrix
145 (e.g., $\Pi_{e_1} = \alpha \frac{\pi_{11}^e}{\tau_{11}} + (1 - \alpha) \frac{\pi_{12}^e}{\tau_{12}}$ for some $0 \leq \alpha \leq 1$). It is the same for the classic bimatrix
146 game except that for us α is no longer a linear function of the strategy frequencies of the
147 other population since the distributional equilibrium is not the standard Hardy–Weinberg
148 distribution. In fact, α depends on population size N as well.

149 A strict NE can be pictured as corresponding to a particular vertex of the unit square
150 (cf. Figure 2 with the axes scaled to be frequencies of the first strategy in each population
151 instead of numbers and with vertices given as solid dots corresponding to strict NE). It is
152 well-known (see Figs. 10.1, 10.2, 11.1 in Hofbauer and Sigmund, 1998, or Figs. 3.3.1, 3.3.2,
153 3.3.3 in Cressman, 2003) that a classic two-strategy bimatrix game may have no strict
154 NE, exactly one strict NE (e.g., Figure 2A), or exactly two strict NE that are diagonally
155 opposite each other (e.g., Figure 2E). Furthermore, the classic two-strategy bimatrix game
156 (with nondegenerate payoff bimatrix) can be classified by its strict NE and its interior
157 NE (i.e., its unique NE where both populations are polymorphic) if it exists.

158 By examining interior NE, we will see this classification method fails for two-strategy
159 time-constrained bimatrix games (see Section 2.2). These equilibria must satisfy $\Pi_{e_1} =$
160 Π_{e_2} and $\Pi_{f_1} = \Pi_{f_2}$ so that neither phenotype can increase its payoff by unilaterally
161 switching its strategy. Unfortunately, obtaining analytic formulas for interior NE seems
162 to be out of reach except in two special cases.

163 One special case is when interaction times satisfy $\tau_{12}\tau_{21} = \tau_{11}\tau_{22}$. Then the payoffs (9)
164 evaluated at the equilibrium distribution (8) are the same as the payoffs for the classic
165 bimatrix game with payoff matrix given by the time adjusted payoff matrix (10), i.e.,

$$\begin{aligned}
\Pi_{e_i} &= \frac{N_{f_1} \pi_{i1}^e}{N \tau_{i1}} + \frac{N_{f_2} \pi_{i2}^e}{N \tau_{i2}} \\
\Pi_{f_j} &= \frac{N_{e_1} \pi_{1j}^f}{N \tau_{1j}} + \frac{N_{e_2} \pi_{2j}^f}{N \tau_{2j}},
\end{aligned} \quad (11)$$

166 where $i, j = 1, 2$, and the interior NE simplifies to

$$(N_{e_1}, N_{f_1}) = \left(\frac{N\tau_{12}(\pi_{22}^f\tau_{21} - \pi_{21}^f\tau_{22})}{\tau_{22}(\pi_{11}^f\tau_{22} - \pi_{12}^f\tau_{21}) + \tau_{12}(\pi_{22}^f\tau_{21} - \pi_{21}^f\tau_{22})}, \frac{N\tau_{21}(\pi_{22}^e\tau_{12} - \pi_{12}^e\tau_{22})}{\tau_{22}(\pi_{11}^e\tau_{22} - \pi_{12}^e\tau_{21}) + \tau_{12}(\pi_{22}^e\tau_{21} - \pi_{21}^e\tau_{22})} \right), \quad (12)$$

167 whenever both components are strictly between 0 and N . In fact, this is the interior NE
168 of the classic bimatrix game with time-adjusted bimatrix (10).

169 The other special case is interior symmetric NE (i.e., those on the main diagonal where
170 $N_{e_1} = N_{f_1}$) for role-independent time constrained bimatrix games. As discussed in Section
171 2.2, there are up to two such diagonal interior symmetric NE and the formulas for these
172 are given in Krivan and Cressman (2017).

173 To find interior NE in the general case, we can instead consider the replicator equation
174 at fixed population size N . This dynamics is given by³

$$\begin{aligned} \frac{dN_{e_1}}{dt} &= \frac{N_{e_1}(N - N_{e_1})}{N} (\Pi_{e_1}(N_{e_1}, N_{f_1}) - \Pi_{e_2}(N_{e_1}, N_{f_1})) \\ \frac{dN_{f_1}}{dt} &= \frac{N_{f_1}(N - N_{f_1})}{N} (\Pi_{f_1}(N_{e_1}, N_{f_1}) - \Pi_{f_2}(N_{e_1}, N_{f_1})) \end{aligned} \quad (13)$$

175 where $\Pi_{e_i}(N_{e_1}, N_{f_1})$ and $\Pi_{f_i}(N_{e_1}, N_{f_1})$ are fitnesses (9) evaluated at the equilibrium dis-
176 tribution (6) for a given (N_{e_1}, N_{f_1}) . Rest points of the replicator equation with N_{e_1} and
177 N_{f_1} strictly between 0 and N are the interior NE of the underlying game (Hofbauer and
178 Sigmund, 1998). Moreover, when all $\tau_{ij} = \tau$ are equal, the dynamics (13) is the replicator
179 equation of the classic bimatrix game (up to the factor τ that only affects the speed along
180 trajectories and not the evolutionary outcome).

181 Through the Owner-Intruder game with time-constraints, we illustrate the two special
182 cases mentioned above (i.e., either $\tau_{12}\tau_{21} = \tau_{11}\tau_{22}$ or interior symmetric NE) as well as
183 the replicator method for the general case.

184 2.2. Owner–Intruder game

185 The classic owner intruder game (Maynard Smith, 1982; Hofbauer and Sigmund, 1998;
186 Cressman, 2003; Broom and Rychtář, 2013) is the two-role extension of the symmetric

³Replicator dynamics at fixed population size assume that frequencies of e_1 strategists p_1 are described by $\frac{dp_1}{dt} = p_1(1 - p_1)(\Pi_{e_1}(N_{e_1}, N_{f_1}) - \Pi_{e_2}(N_{e_1}, N_{f_1}))$ (Hofbauer and Sigmund, 1998). Because $N_{e_1} = p_1N$ and the overall size N of population 1 is assumed to be fixed, we obtain $\frac{dN_{e_1}}{dt} = \frac{dp_1}{dt}N$ which yields the first equation in (13).

187 Hawk–Dove game (i.e., matrix game) that models the situation in which an individual
 188 either owns a site or is an intruder trying to seize a site. An individual can either be a
 189 Hawk (strategy e_1 if owner and f_1 if intruder) or a Dove (strategy e_2 if owner and f_2 if
 190 intruder) in either of the two roles. The payoff bimatrix of the game is

$$\begin{array}{c|cc} \text{Owner}\backslash\text{Intruder} & \text{Hawk} & \text{Dove} \\ \hline \text{Hawk} & \left[\frac{V-C}{2}, \frac{V-C}{2} \right] & [V, 0] \\ \text{Dove} & [0, V] & \left[\frac{V}{2}, \frac{V}{2} \right] \end{array}$$

191 where V (the value attached to the site) and C (the cost of fighting) are positive. It is an
 192 example of a role-independent bimatrix game since an individual’s payoff depends only
 193 on the strategies used in the interaction and not on whether the individual is the owner
 194 or the intruder.⁴

195 When the cost of fighting is low ($C < V$), the classic game has a single NE $(e_1, f_1) =$
 196 (H, H) where individuals in both positions behave as hawks. When the cost of fighting is
 197 high ($C > V$) there are two strict NE $(e_2, f_1) = (D, H)$ and $(e_1, f_2) = (H, D)$ as well as a
 198 mixed NE $(p_1, q_1) = (V/C, V/C)$, where Hawk strategy is played with probability V/C in
 199 both roles. This mixed NE cannot be a (two-species) ESS, because bimatrix games can
 200 have ESSs only in pure strategies (Selten, 1980).⁵

201 For the time-constrained bimatrix game, we first analyze its strict NE through the
 202 following time-adjusted bimatrix (cf. (10))

$$\begin{array}{c|cc} \text{Owner}\backslash\text{Intruder} & \text{Hawk} & \text{Dove} \\ \hline \text{Hawk} & \left[\frac{V-C}{2\tau_{11}}, \frac{V-C}{2\tau_{11}} \right] & \left[\frac{V}{\tau_{12}}, 0 \right] \\ \text{Dove} & \left[0, \frac{V}{\tau_{21}} \right] & \left[\frac{V}{2\tau_{22}}, \frac{V}{2\tau_{22}} \right] \end{array}.$$

203 The following list contains all strict NE of the time-constrained Owner–Intruder game
 204 (Figure 1). After each item in this list, the panels in Figure 2 that have this strict NE
 205 are indicated in parentheses.

- 206 • If $V > C$, then strategy (H, H) is a NE (e.g., Figure 2A–D).

⁴Broom and Rychtář (2013) refer to role independence as an “uncorrelated asymmetry” (see also the role games of Hofbauer and Sigmund (1998)). Mathematically, role independence is equivalent to the second payoff entries in the bimatrix forming the transpose of the matrix of first entries. It is assumed that the pure strategy sets for both roles are the same as well as the ordering of their elements. Typically, the strategies are given the same name in both roles (e.g., Hawk and Dove) and the same order. Every role-independent bimatrix game is the two-role extension of a symmetric matrix game and has NE where both populations use the same strategy; namely, a NE of the matrix game. In addition, there may be other NE.

⁵In fact, a strategy pair is an ESS for a classic bimatrix game if and only if it is a strict NE.

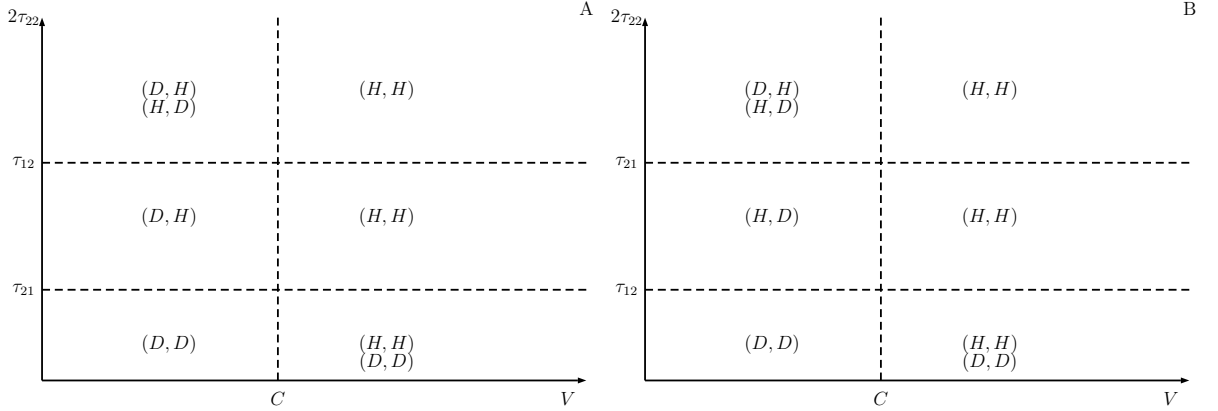


Figure 1: Strict NE of the Owner–Intruder game as functions of V and $2\tau_{22}$ parameters. Panel A assumes that $\tau_{21} < \tau_{12}$ and panel B assumes the opposite inequality.

- 207 • If $\tau_{12} > 2\tau_{22}$ and $\tau_{21} > 2\tau_{22}$, then strategy (D, D) is a NE (e.g., Figure 2B, F, G,
208 H).
- 209 • If $V < C$ and $\tau_{12} < 2\tau_{22}$, then strategy (H, D) is a NE (e.g., Figure 2E, J).
- 210 • If $V < C$ and $\tau_{21} < 2\tau_{22}$, then strategy (D, H) is a NE (e.g., Figure 2E, I).

211 Dependence of strict NEs as a function of model parameters are shown in Figure 1. There
212 is at least one strict NE for all parameter values except in the degenerate situations where
213 $V = C$, $\tau_{12} = 2\tau_{22}$, or $\tau_{21} = 2\tau_{22}$ (these are the dashed lines in Figure 1) that are discussed
214 in Section 2.3.

215 Of particular note is that, although strategy pair (Dove, Dove) is never an ESS (i.e.
216 a strict NE) for the classic Owner–Intruder game (since Dove is never an ESS for the
217 Hawk–Dove matrix game), this pair is a strict NE when $2\tau_{22} < \min\{\tau_{12}, \tau_{21}\}$. This analysis
218 shows that when compared with the classical model, the model that considers duration
219 of interactions can have strategy (D, D) as a NE provided the interaction time between
220 Doves is small.

In the special case where interaction times satisfy $\tau_{12}\tau_{21} = \tau_{11}\tau_{22}$, the interior NE (provided it exists) is given by (12) as

$$(N_{e1}, N_{f1}) = \left(\frac{N\tau_{12}V(\tau_{21} - 2\tau_{22})}{\tau_{22}^2(V - C) + \tau_{12}V(\tau_{21} - 2\tau_{22})}, \frac{N\tau_{21}V(\tau_{12} - 2\tau_{22})}{(V - C)\tau_{22}^2 + \tau_{21}V(\tau_{12} - 2\tau_{22})} \right).$$

221 We observe that when all interaction times are the same, the interior equilibrium is
222 $(N_{e1}, N_{f1}) = (V/C, V/C)$ exactly as in the classical Owner–Intruder game.

223 To investigate interior NE further for the Owner–Intruder game, fitness functions (9)
 224 are now

$$\begin{aligned}
 \Pi_{e_1} &= \frac{n_{11}(V - C)}{2\tau_{11}(n_{11} + n_{12})} + \frac{n_{12}V}{\tau_{12}(n_{11} + n_{12})}, \\
 \Pi_{e_2} &= \frac{n_{22}V}{2\tau_{22}(n_{21} + n_{22})}, \\
 \Pi_{f_1} &= \frac{n_{11}(V - C)}{2\tau_{11}(n_{11} + n_{21})} + \frac{n_{21}V}{\tau_{21}(n_{11} + n_{21})}, \\
 \Pi_{f_2} &= \frac{n_{22}V}{2\tau_{22}(n_{12} + n_{22})}.
 \end{aligned} \tag{14}$$

225 Evaluating these at the equilibrium distribution (6) yields

$$\begin{aligned}
 \Pi_{e_1} &= \frac{(C\tau_{12} + (2\tau_{11} - \tau_{12})V)(\sqrt{A} - N\tau_{12}\tau_{21})}{4N_{e_1}\tau_{11}\tau_{12}(\tau_{11}\tau_{22} - \tau_{12}\tau_{21})} + \frac{\tau_{12}(V - C)(N_{e_1} + N_{f_1}) + 2\tau_{11}V(N_{e_1} - N_{f_1})}{4N_{e_1}\tau_{11}\tau_{12}} \\
 \Pi_{e_2} &= - \frac{V \left(\sqrt{A} + N(\tau_{12}\tau_{21} - 2\tau_{11}\tau_{22}) - (N_{e_1} + N_{f_1})(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}) \right)}{4\tau_{22}(N - N_{e_1})(\tau_{11}\tau_{22} - \tau_{12}\tau_{21})} \\
 \Pi_{f_1} &= - \frac{\sqrt{A}(C\tau_{21} + (2\tau_{11} - \tau_{21})V)}{4N_{f_1}\tau_{11}\tau_{21}(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})} + \frac{N\tau_{12}(C\tau_{21} + 2\tau_{11}V - \tau_{21}V)}{4N_{f_1}\tau_{11}(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})} - \\
 &\quad \frac{(C - V)(N_{e_1} + N_{f_1})}{4N_{f_1}\tau_{11}} + \frac{2\tau_{11}V(N_{f_1} - N_{e_1})}{4N_{f_1}\tau_{11}\tau_{21}} \\
 \Pi_{f_2} &= - \frac{V \left(\sqrt{A} + N(\tau_{12}\tau_{21} - 2\tau_{11}\tau_{22}) - (N_{e_1} + N_{f_1})(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}) \right)}{4\tau_{22}(N - N_{f_1})(\tau_{11}\tau_{22} - \tau_{12}\tau_{21})},
 \end{aligned}$$

226 where A is given in (7). To find interior NE, we need to solve $\Pi_{e_1} = \Pi_{e_2}$ and $\Pi_{f_1} = \Pi_{f_2}$.

227 Two-strategy, bimatrix games that are role-independent have role-independent inter-
 228 action times if and only if $\tau_{12} = \tau_{21}$ (i.e., the length of Hawk–Dove interactions does
 229 not depend on whether the Hawk is the owner or the intruder).⁶ Symmetric NE of
 230 the role-independent time-constrained Owner–Intruder game are then those of the time-
 231 constrained Hawk–Dove matrix game, which are found analytically in Křivan and Cress-
 232 man (2017) using `Solve` command of Mathematica 11.

233 Since attempts to use this method to find interior NE when the time-constrained
 234 bimatrix game was not role-independent or interaction times did not satisfy $\tau_{12}\tau_{21} = \tau_{11}\tau_{22}$

⁶We call a multi-strategy time-constrained bimatrix game “role-independent” if both its payoff bima-
 trix and its time interaction matrix are role-independent. This last requirement is equivalent to the time
 interaction matrix being symmetric (i.e., $\tau_{ij} = \tau_{ji}$ for all i, j).

235 failed, we now analyze the NE of the Owner–Intruder game numerically through the
236 replicator equation, focusing on the cases where $V > C$ and $V < C$ separately.

237 First, assume that $V > C$ (Figure 2, panels A–D). Then (H, H) is always a strict
238 NE. When the time-constrained Owner–Intruder game is role-independent, the replicator
239 equation is invariant along the main diagonal of the unit square and its trajectories in
240 the unit square are reflections in the main diagonal (Figure 2A,B,C). Furthermore, on the
241 diagonal, the dynamics (13) restricts to the replicator equation for the time-constrained
242 Hawk–Dove matrix game, which was analyzed by Křivan and Cressman (2017). They
243 showed that, when interaction times between two Hawks are long enough (and all other
244 interaction times are the same), there exist two (symmetric) interior NEs and the one with
245 fewer Hawks is locally asymptotically stable while the other one is unstable. However,
246 numerical simulations (e.g., Figure 2C) show that both interior symmetric NE (i.e., those
247 gray points that are on the main diagonal) are saddles (i.e., unstable) for the bimatrix
248 replicator dynamics.⁷

249 Simulations of the replicator equation for the role-independent time-constrained Owner–
250 Intruder game with $V > C$ show that long interaction times between Hawks now lead to
251 two new asymmetric interior NE (i.e., those off the main diagonal shown as black interior
252 dots in Figure 2C). Numerical simulations suggest that these two equilibria are neutrally
253 stable as they appear to be surrounded by a family of closed trajectories. The domain of
254 the phase space filled by these closed curves is separated from the rest by two heteroclinic
255 orbits that join the two symmetric NE. In particular, the symmetric strict NE (H, H)
256 where all individuals play Hawk is not globally asymptotically stable.

257 The neutral stability of the asymmetric NE disappears when the time interaction
258 matrix is role dependent. For example, it is reasonable to assume that interaction time
259 between intruding Hawk and owning Dove is longer than that between intruding Dove
260 and owning Hawk (i.e., $\tau_{21} > \tau_{12}$) because an owning Dove tries to defend its site against
261 attacking Hawk. This role-dependent interaction time makes one of the two interior
262 asymmetric NE unstable while the other becomes locally asymptotically stable (Figure
263 2D).

264 Now assume that $V < C$ (Figure 2, panels E–K). Hawk is no longer an ESS for

⁷From extensive simulations of the replicator equation, it seems likely that any interior symmetric NE of two-strategy role-independent time-constrained bimatrix games are always saddles but we have no proof of this conjecture. In the special case where $\tau_{12}\tau_{21} = \tau_{11}\tau_{22}$ (and $\tau_{12} = \tau_{21}$), interior symmetric NE are saddles since, from (11), Π_{e_1} (and Π_{f_1}) depends only on the strategy frequency of the other population, implying that the Jacobian of replicator dynamics (13) evaluated at interior equilibrium (12) has zeros on the main diagonal. This extends the same well-known result for classic role-independent bimatrix games (Hofbauer and Sigmund, 1998).

265 the classic Hawk–Dove game and the only NE is the interior ESS where the population
 266 plays Hawk with probability $\frac{V}{C}$. On the other hand, the classic Owner–Intruder game has
 267 two strict NE (H, D) and (D, H) ⁸ and the unstable interior NE where both populations
 268 plays Hawk with probability $\frac{V}{C}$. This corresponds to the time-constrained game with all
 269 interaction times equal (Figure 2E). When Hawk–Dove interactions are sufficiently long
 270 compared to Dove–Dove interactions (specifically, $\tau_{21} > 2\tau_{22}$ and $\tau_{12} > 2\tau_{22}$), then (D, D)
 271 is the only NE (Figure 2F). With a lower cost (Figure 2G), two neutrally stable symmetric
 272 interior NE that are surrounded by a family of closed trajectories appear. Furthermore,
 273 a small perturbation of these NE by introducing a slight role dependence in interaction
 274 times makes one of them locally asymptotically stable and the other unstable (panel
 275 H). Larger differences for role dependent interaction times (panels I and J respectively)
 276 eliminates interior NE altogether and make the paradoxical ESS (D, H) (respectively,
 277 (H, D)) globally asymptotically stable. Panel K is a degenerate case where $\tau_{12} = \tau_{21} =$
 278 $2\tau_{22}$ and so has boundary NE as discussed in the following section. Finally, panel L
 279 assumes $V = C = 1$, $\tau_{11} = 3$ and all other interaction times are 1. This parametrization
 280 corresponds to the situation where sets of the NE along the boundary of square $[0, N] \times$
 281 $[0, N]$ exist. As calculated in the following section, the sets of NE are $0 \leq N_{e_1} < \frac{3}{4}N$
 282 when $N_{f_1} = N$ and $0 \leq N_{f_1} < \frac{3}{4}N$ when $N_{e_1} = N$.

283 2.3. Boundary NE

284 The previous two sections analyzed the strict NE and interior NE for two-strategy
 285 time-constrained bimatrix games. These games may also have NE on an edge of the
 286 square that are not at a vertex (i.e., partially mixed NE where only one of the two
 287 populations is polymorphic). For example, suppose that population 1 is polymorphic
 288 and population 2 is monomorphic at pure strategy f_1 , i.e., $N_{f_1} = N$. Then, at a NE on
 289 this edge, the fitnesses of both strategies of population 1 must be equal, i.e., $\Pi_{e_1} = \Pi_{e_2}$.
 290 Since $n_{12} = n_{22} = 0$, $\Pi_{e_1} = \frac{\pi_{11}^e}{\tau_{11}}$ and $\Pi_{e_2} = \frac{\pi_{21}^e}{\tau_{21}}$ from (9).⁹ In this degenerate case where
 291 $\frac{\pi_{11}^e}{\tau_{11}} = \frac{\pi_{21}^e}{\tau_{21}}$, a point along the edge $N_{f_1} = N$ is a NE if and only if $\Pi_{f_1} \geq \Pi_{f_2}$. Since
 292 $n_{21} = N_{e_2}$, $n_{11} = N_{e_1}$ and $N = N_{e_1} + N_{e_2}$,

$$\Pi_{f_1} = \frac{N_{e_1}}{N} \frac{\pi_{11}^f}{\tau_{11}} + \frac{N_{e_2}}{N} \frac{\pi_{21}^f}{\tau_{21}}. \quad (15)$$

⁸The second strict NE is often called the “paradoxical ESS” (Maynard Smith, 1982) since it corresponds to the intruder always taking over the site and becoming the owner.

⁹In classic two-strategy bimatrix games, the pure strategy pair (e_1, f_1) may be a NE in this situation but not a strict NE. We have ignored this degenerate case in the classification of pure strategy NE in Sections 2.1 and 2.2 of our time-constrained bimatrix game through (10) above.

293 On the other hand, the invasion fitness of strategy f_2 when there are no individuals playing
 294 this strategy is (see Appendix B)

$$\Pi_{f_2} = \frac{N_{e_1} \pi_{12}^f \tau_{21} + N_{e_2} \pi_{22}^f \tau_{11}}{N \tau_{11} \tau_{22} + N_{e_1} (\tau_{12} \tau_{21} - \tau_{11} \tau_{22})}. \quad (16)$$

295 Solving $\Pi_{f_1} = \Pi_{f_2}$ gives us, in general, up to two roots for N_{e_1} satisfying $0 \leq N_{e_1} \leq N$.
 296 These roots divide the edge into closed subintervals, on each of which $\Pi_{f_1} - \Pi_{f_2}$ does not
 297 change sign. Each such subinterval with this difference nonnegative is then a connected set
 298 of NE.¹⁰ However, since each point on this edge is a rest point of the replicator equation,
 299 none can be asymptotically stable under this dynamics.

300 For the Owner–Intruder game, boundary NE emerge on the top edge of the square
 301 $[0, N] \times [0, N]$ where $N_{f_1} = N$ when $V = C$ since $\frac{\pi_{11}^e}{\tau_{11}} = \frac{\pi_{21}^e}{\tau_{21}} = 0$ along this edge. By
 302 evaluating when $\Pi_{f_1} \geq \Pi_{f_2}$ along this edge, we find the following four cases for sets of NE
 303 of the form (N_{e_1}, N)

- 304 1. $\tau_{11} \leq 2\tau_{12}$ and $\tau_{21} < 2\tau_{22}$ and $0 \leq N_{e_1} \leq N$
- 305 2. $\tau_{11} > 2\tau_{12}$ and $\tau_{21} < 2\tau_{22}$ and $0 \leq N_{e_1} \leq \frac{N\tau_{11}(\tau_{21}-2\tau_{22})}{2(\tau_{12}\tau_{21}-\tau_{11}\tau_{22})}$
- 306 3. $\tau_{11} < 2\tau_{12}$ and $\tau_{21} \geq 2\tau_{22}$ and $\frac{N\tau_{11}(\tau_{21}-2\tau_{22})}{2(\tau_{12}\tau_{21}-\tau_{11}\tau_{22})} \leq N_{e_1} \leq N$

307 Similarly, let us consider the right edge of the square where all individuals of the first
 308 species play strategy Hawk, i.e., $N_{e_1} = N$. When $V = C$, this leads to the following sets
 309 of NE for the Owner–Intruder game.

- 310 1. $\tau_{11} \leq 2\tau_{21}$ and $\tau_{12} < 2\tau_{22}$ and $0 \leq N_{f_1} \leq N$
- 311 2. $\tau_{11} > 2\tau_{21}$ and $\tau_{12} < 2\tau_{22}$ and $0 \leq N_{f_1} \leq \frac{N\tau_{11}(\tau_{12}-2\tau_{22})}{2(\tau_{12}\tau_{21}-\tau_{11}\tau_{22})}$
- 312 3. $\tau_{11} < 2\tau_{21}$ and $\tau_{12} > 2\tau_{22}$ and $\frac{N\tau_{11}(\tau_{12}-2\tau_{22})}{2(\tau_{12}\tau_{21}-\tau_{11}\tau_{22})} \leq N_{f_1} \leq N$

313 These sets of NE on the boundary are illustrated in Figure 2L for the role-independent
 314 time-constrained Owner–Intruder game with $V = C$. From Krivan and Cressman (2017)
 315 the interior NE in this figure appears for $\tau_{11} > \tau(3 - C/V + 2\sqrt{1 - C/V}) = 2\tau$ (assuming
 316 $\tau_{12} = \tau_{22}$). In this case the NEs on the edges form two disconnected components. Since
 317 $\tau_{12} = \tau_{21}$, the NE component on the upper edge is then the reflection in the main diagonal
 318 of the component on the right-hand edge.

319 We note that sets of NE also appear (Figure 2K) on the lower (respectively, left-
 320 hand) edges of the square when $\tau_{12} = 2\tau_{22}$ (respectively, $\tau_{21} = 2\tau_{22}$). By other choices of
 321 interaction time τ_{11} we can also get disconnected components along these edges.

¹⁰In classical games, this set is called a NE component (Cressman, 2003).

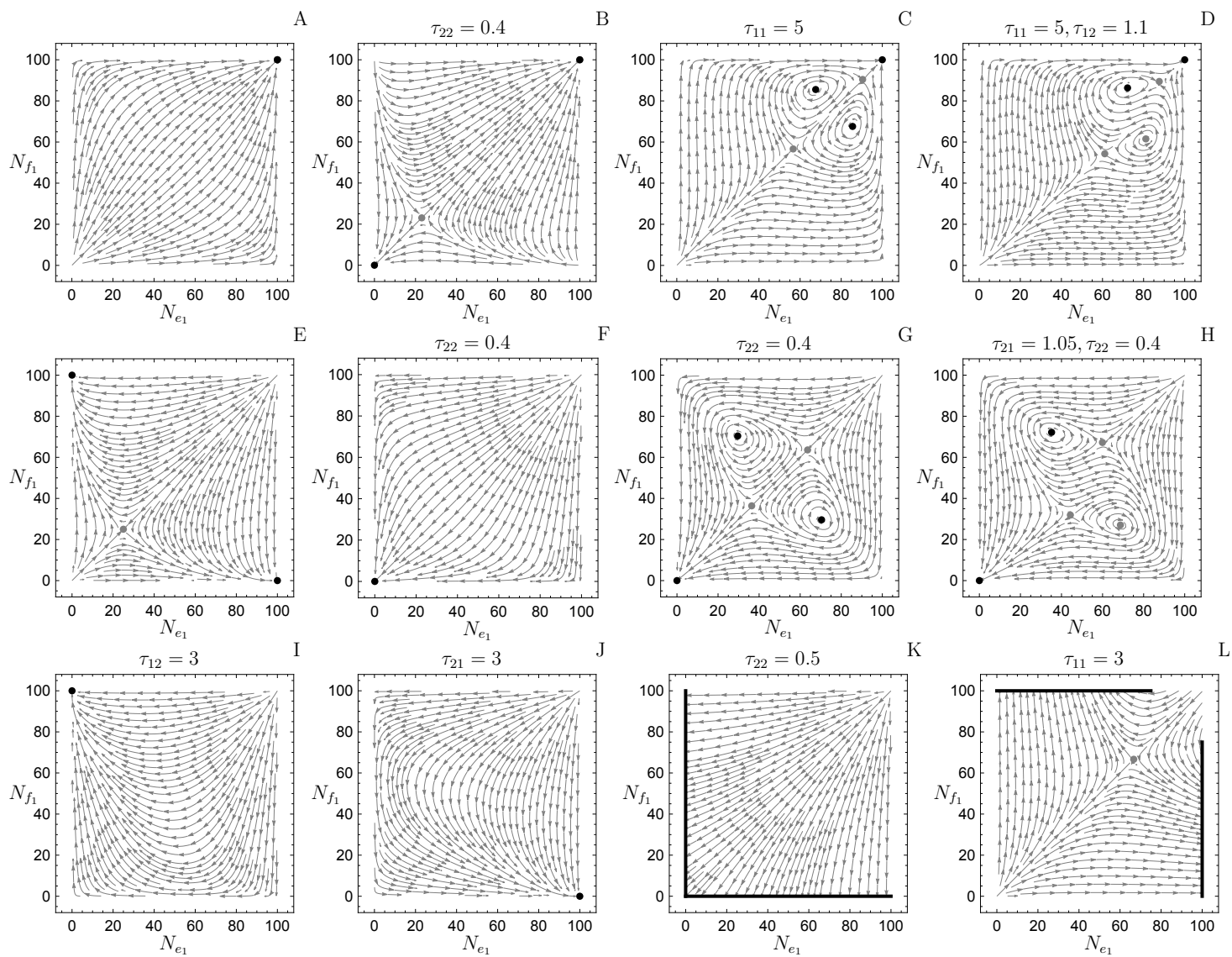


Figure 2: The replicator dynamics for the Owner–Intruder game depending on V , C and interaction times when pairing is instantaneous. The first four panels (i.e., panels A, B, C, and D) assume $V > C$ (in fact, $V = 4$ and $C = 1$). The other panels assume $V = 1 \leq C$ with $C = 4$ (panels E, F, I, J, K), $C = 1.5$ (panels G, H) and $C = 1$ (panel L). All interaction times not equal to 1 are indicated in each panel. Thus, panels A and E respectively are the replicator dynamics of the classic Owner–Intruder game for $V > C$ and $V < C$ respectively since all interaction times are the same. In particular, the main diagonal is invariant in these two panels since the time-constrained game is role-independent. For the same reason, this invariance holds in panels B, C, F, G, K, L but not in the other four panels (D, H, I, J) that have role dependent interaction times (i.e., $\tau_{12} \neq \tau_{21}$). In panel B, strategy pairs (H, H) and (D, D) are strict NE (since $\min\{\tau_{12}, \tau_{21}\} > 2\tau_{22}$ and $V > C$) and an unstable saddle symmetric interior NE appears. In panel C, Hawk-Hawk interaction time is long enough ($\tau_{11} = 5$) that two unstable saddle symmetric interior NE emerge along with two neutrally stable asymmetric ones. Panel D is an asymmetric perturbation of the interaction time matrix from panel C (specifically τ_{12} shifts from 1 to 1.1) that perturbs the two asymmetric NE to a stable and unstable one. Since $\min\{\tau_{12}, \tau_{21}\} > 2\tau_{22}$ and $V < C$ in panels F, G, H, (D, D) is the only strict NE. It may be globally asymptotically stable (panel F) or only locally asymptotically stable when there are four interior NE with two unstable saddles and two neutrally stable (the role-independent case of panel G) or two unstable saddles together with one unstable and one stable NE (panel H with perturbed interaction matrix compared to panel G). In the role-dependent interaction matrices of panels I and J, τ_{12} (respectively τ_{21}) is large enough that the paradoxical ESS (D, H) (respectively (H, D)) is the only strict NE and it is globally asymptotically stable. Finally, panels K and L illustrate that sets of boundary NE emerge (thick black line segments) when $V = 1$, $C = 4$, $\tau_{11} = \tau_{12} = \tau_{21} = 2\tau_{22} = 1$ (panel K), and $V = C$ (panel L).

322 3. Non instantaneous pair formation

323 So far we have assumed that pair formation is instantaneous, i.e., there are no singles.
 324 This assumption is natural in population genetics, where alleles exist as singles only during
 325 meiosis but otherwise they are always paired in diploid individuals. However, since it may
 326 be more realistic in general to assume that it takes some time for singles to form pairs, we
 327 consider both singles and paired individuals in this section. We also assume that, when a
 328 pair disbands, these new singles are ready immediately to start searching for new partners
 329 with encounter rate λ and new pairs are formed by random encounters between one single
 330 from each population.¹¹

¹¹These last assumptions rule out applying the methods to bimatrix games where newly single individuals may wait after disbanding before they are ready to form new pairs. For example, in the model

331 The number of singles of the two strategies for population 1 are denoted by n_{e_i} for
 332 $i = 1, 2$ and for population 2 by n_{f_j} for $j = 1, 2$. Then

$$\begin{aligned} N_{e_i} &= n_{e_i} + n_{i1} + n_{i2} \\ N_{f_j} &= n_{f_j} + n_{1j} + n_{2j} \end{aligned} \tag{17}$$

333 are the total number of individuals playing a given strategy. We continue to assume that
 334 the total number of individuals in each population is N (i.e., $N_{e_1} + N_{e_2} = N = N_{f_1} + N_{f_2}$).

335 Distributional dynamics of singles and pairs when pair formation is described by the
 336 mass action law are then

$$\begin{aligned} \frac{dn_{e_1}}{dt} &= -\lambda n_{e_1}(n_{f_1} + n_{f_2}) + \frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}} \\ \frac{dn_{e_2}}{dt} &= -\lambda n_{e_2}(n_{f_1} + n_{f_2}) + \frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}} \\ \frac{dn_{f_1}}{dt} &= -\lambda n_{f_1}(n_{e_1} + n_{e_2}) + \frac{n_{11}}{\tau_{11}} + \frac{n_{21}}{\tau_{21}} \\ \frac{dn_{f_2}}{dt} &= -\lambda n_{f_2}(n_{e_1} + n_{e_2}) + \frac{n_{12}}{\tau_{12}} + \frac{n_{22}}{\tau_{22}} \\ \frac{dn_{11}}{dt} &= \lambda n_{e_1} n_{f_1} - \frac{n_{11}}{\tau_{11}} \\ \frac{dn_{12}}{dt} &= \lambda n_{e_1} n_{f_2} - \frac{n_{12}}{\tau_{12}} \\ \frac{dn_{21}}{dt} &= \lambda n_{e_2} n_{f_1} - \frac{n_{21}}{\tau_{21}} \\ \frac{dn_{22}}{dt} &= \lambda n_{e_2} n_{f_2} - \frac{n_{22}}{\tau_{22}}. \end{aligned} \tag{18}$$

337 Appendix C shows that (18) has a unique distributional equilibrium for a fixed N and
 338 given N_{e_1} and N_{f_1} .

339 Assuming that singles do not get any payoffs, the fitnesses (i.e., the expected payoff
 340 to an individual per unit time) of the four strategies evaluated at the unique equilibrium

for parental care of offspring known as the Battle of the Sexes (Dawkins, 1976), when fast females mate with philandering males to produce offspring, it is assumed that the male immediately deserts and begins searching for a new mate whereas the female remains and cares for the offspring for a certain amount of time before searching for a new mate.

341 of (18) are $(i, j = 1, 2)$

$$\begin{aligned}\Pi_{e_i} &= \frac{n_{i1}}{N_{e_i}} \frac{\pi_{i1}^e}{\tau_{i1}} + \frac{n_{i2}}{N_{e_i}} \frac{\pi_{i2}^e}{\tau_{i2}}, \\ \Pi_{f_j} &= \frac{n_{1j}}{N_{f_j}} \frac{\pi_{1j}^f}{\tau_{1j}} + \frac{n_{2j}}{N_{f_j}} \frac{\pi_{2j}^f}{\tau_{2j}}.\end{aligned}\tag{19}$$

342 These fitness functions depend on N , N_{e_1} and N_{f_1} . Since, at the unique distributional
343 equilibrium of (18),

$$n_{ij} = \lambda n_{e_i} n_{f_i} \tau_{ij}, \quad i, j = 1, 2\tag{20}$$

344 fitnesses (19) simplify to $(i, j = 1, 2)$

$$\begin{aligned}\Pi_{e_i} &= \frac{\lambda(n_{f_1} \pi_{i1}^e + n_{f_2} \pi_{i2}^e)}{\lambda n_{f_1} \tau_{i1} + \lambda n_{f_2} \tau_{i2} + 1}, \\ \Pi_{f_j} &= \frac{\lambda(n_{e_1} \pi_{1j}^f + n_{e_2} \pi_{2j}^f)}{\lambda n_{e_1} \tau_{1j} + \lambda n_{e_2} \tau_{2j} + 1}.\end{aligned}\tag{21}$$

345 The time-constrained bimatrix game with non instantaneous pair formation based on
346 payoff bimatrix (1) and time interaction matrix (2) is then the two-strategy game with
347 payoffs given by the fitness functions (21) evaluated at the distributional equilibrium of
348 (18) for fixed size N of each population and encounter rate λ . As in Section 2, we are
349 interested in the NE of this game and its evolutionary outcome.

350 3.1. Classic bimatrix game with non instantaneous pair formation

351 The classic model implicitly assumes all interaction times are equal (i.e., $\tau_{ij} = \tau$ for
352 all $i, j = 1, 2$). However, since the classic model also assumes that individuals are always
353 interacting (i.e., always in pairs), the question arises whether the classic predictions remain
354 valid when pair formation requires time. This section examines the question.

355 The equilibrium distribution of (18) is

$$\begin{aligned}n_{e_i} &= \frac{N_{e_i}}{N} \frac{(\sqrt{4\lambda N\tau} + 1 - 1)}{2\lambda N\tau} \\ n_{f_j} &= \frac{N_{f_j}}{N} \frac{(\sqrt{4\lambda N\tau} + 1 - 1)}{2\lambda N\tau}.\end{aligned}$$

356 Substituting these expressions to (21) leads to

$$\begin{aligned}\Pi_{e_i} &= \frac{4N\lambda}{(1 + \sqrt{1 + 4\lambda\tau N})^2} \left(\pi_{i1}^e \frac{N_{f_1}}{N} + \pi_{i2}^e \frac{N_{f_2}}{N} \right), \\ \Pi_{f_j} &= \frac{4N\lambda}{(1 + \sqrt{1 + 4\lambda\tau N})^2} \left(\pi_{1j}^f \frac{N_{e_1}}{N} + \pi_{2j}^f \frac{N_{e_2}}{N} \right).\end{aligned}\tag{22}$$

357 Thus, up to the positive factor $\frac{4N\lambda}{(1+\sqrt{1+4\lambda\tau N})^2}$, these are the payoffs of the classic bimatrix
 358 game with payoff matrix (1). From this it follows that the NE of the classic bimatrix
 359 game with non instantaneous pair formation is the same as the the classic bimatrix game
 360 and, moreover, the trajectories of the replicator equation are the same (up to the speed
 361 along the trajectory). Thus, the two games have the same evolutionary outcomes.

362 To rephrase, standard evolutionary game theory models of bimatrix games can ex-
 363 plicitly incorporate time constraints without affecting the game-theoretic analysis as long
 364 as all interaction times are the same. It is then irrelevant whether pair formation is
 365 instantaneous or requires some time.

366 3.2. Evolutionary outcomes with non instantaneous pair formation

367 As we saw in Section 2, evolutionary outcomes of time-constrained bimatrix games
 368 with instantaneous pair formation depend heavily on pair interaction times when these
 369 are not all the same (e.g., Figure 2). This section analyzes the same phenomena when
 370 pair formation is not instantaneous.

371 We start by characterizing the strict NE of these games. From (21), at strategy pair
 372 (e_1, f_1) ,

$$\Pi_{e_1} = \frac{\lambda n_{f_1} \pi_{11}^e}{\lambda n_{f_1} \tau_{11} + 1}, \quad \Pi_{f_1} = \frac{\lambda n_{e_1} \pi_{11}^f}{\lambda n_{e_1} \tau_{11} + 1},$$

373 since $n_{e_2} = n_{f_2} = 0$. Note that the fitness Π_{e_1} (Π_{f_1}) does not depend on distributional
 374 equilibrium of population 1 (2). Thus, the invasion fitnesses of strategy e_2 and f_2 are

$$\Pi_{e_2} = \frac{\lambda n_{f_1} \pi_{21}^e}{\lambda n_{f_1} \tau_{21} + 1}$$

375 and

$$\Pi_{f_2} = \frac{\lambda n_{e_1} \pi_{12}^f}{\lambda n_{e_1} \tau_{12} + 1}$$

376 as given in (21) with $n_{e_2} = n_{f_2} = 0$. Furthermore, at this strategy pair, $N = n_{e_1} + n_{11} =$
 377 $n_{e_1} + \lambda n_{e_1} n_{f_1} \tau_{11} = n_{f_1} + n_{11}$. Thus, $n_{e_1} = n_{f_1}$ and so $N = \lambda \tau_{11} n_{e_1}^2 + n_{e_1}$ and

$$n_{e_1} = \frac{-1 + \sqrt{1 + 4N\lambda\tau_{11}}}{2\lambda\tau_{11}} = n_{f_1}.$$

378 Strategy pair (e_1, f_1) is a strict NE provided $\Pi_{e_1} > \Pi_{e_2}$ and $\Pi_{f_1} > \Pi_{f_2}$, i.e.,

$$\begin{aligned} \frac{\pi_{11}^e}{\tau_{11}(\sqrt{4\lambda N\tau_{11} + 1} + 1)} &> \frac{\pi_{21}^e}{\tau_{21}(\sqrt{4\lambda N\tau_{11} + 1} - 1) + 2\tau_{11}} \\ \frac{\pi_{11}^f}{\tau_{11}(\sqrt{4\lambda N\tau_{11} + 1} + 1)} &> \frac{\pi_{12}^f}{\tau_{12}(\sqrt{4\lambda N\tau_{11} + 1} - 1) + 2\tau_{11}}. \end{aligned} \tag{23}$$

379 Similarly, we can obtain conditions for other strict NE. Contrary to the case of instan-
 380 tantaneous pairing where these conditions are given by the adjusted payoff matrix (10), we
 381 cannot write these conditions in a similar form when pairing is non-instantaneous. This
 382 is seen from expressions (23), where the invasion fitness for strategy e_2 (f_2) depends not
 383 only on interaction time τ_{21} (τ_{12}), but also on interaction time τ_{11} .

384 As λ increases to infinity, payoff Π_{e_1} (Π_{f_1}) converges to π_{11}^e/τ_{11} (π_{11}^f/τ_{11}) and invasion
 385 fitness Π_{e_2} (Π_{f_2}) converges to π_{21}^e/τ_{21} (π_{12}^f/τ_{12}). Thus, when the encounter rate of singles
 386 is large, the strict NE of the time-constrained bimatrix game with non instantaneous pair
 387 formation are the same as the strict NE of the time-constrained bimatrix game of Section
 388 2 (i.e., with instantaneous pair formation). In fact, for large λ , the interior NE match as
 389 well since there are essentially no singles in the system.

390 The next section illustrates these general results for the Owner–Intruder game.

391 3.3. The Owner–Intruder game with non instantaneous pair formation

392 When all interaction times equal to τ as in Section 3.1, there is an interior NE if and
 393 only if $V < C$. As a function of λ and τ , it is given by

$$\begin{aligned}
 n_{e_1} = n_{f_1} &= \frac{V(-1 + \sqrt{1 + 4\lambda N\tau})}{2C\lambda\tau} \\
 n_{e_2} = n_{f_2} &= \frac{(C - V)(-1 + \sqrt{4\lambda N\tau})}{2C\lambda\tau} \\
 N_{e_1} = N_{f_1} &= \frac{NV}{C},
 \end{aligned}
 \tag{24}$$

394 which is the classic result for the case when $V < C$.

395 However, for a general time interaction matrix, an analytic expression for the interior
 396 NE is not available. Our recourse is to apply the replicator equation (13) with payoffs
 397 (21) when pairing is non-instantaneous. On contrary to the case of instantaneous pairing,
 398 we cannot now express the distributional equilibrium at the current strategy numbers
 399 explicitly. Thus, we have to solve replicator equation (13) together with the system of
 400 algebraic equations

$$\begin{aligned}
 N_{e_1} &= n_{e_1}(1 + \lambda n_{f_1}\tau_{11} + \lambda n_{f_2}\tau_{12}) \\
 N_{f_1} &= n_{f_1}(1 + \lambda n_{e_1}\tau_{11} + \lambda n_{e_2}\tau_{21}).
 \end{aligned}
 \tag{25}$$

401 This is a semi-explicit index 1 differential-algebraic equation (Ascher and Petzold, 1998)
 402 that we solve numerically using Mathematica 10.

403 Figure 3 shows the results for two encounter rates. Panels A–H use the same parameter
 404 values (i.e., V , C , τ_{ij}) as corresponding panels in Figure 2. For the role-independent time-
 405 constrained cases (panels A–C, E–G), trajectories remain reflections of each other with

406 respect to the main diagonal. We see that for large enough encounter rate ($\lambda = 10$ in
407 panels A–H) the strict NE still match those of Section 2. However, there are differences
408 in stability of interior NE between Figures 2 and 3. The neutral stability of the two
409 off-diagonal equilibria in Figure 2C and G is lost and the two equilibria become unstable.
410 Figure 3C and G show two trajectories that start close to the two equilibria and that
411 converge to equilibrium $(N_{e_1}, N_{f_1}) = (100, 100)$ and $(N_{e_1}, N_{f_1}) = (0, 0)$, respectively.
412 Panels C', D', G', and H' show numerical simulations for yet smaller encounter rate
413 ($\lambda = 1$). We observe that this leads to disappearance of interior NE in panels C' and D',
414 and to destabilization of the interior stable NE in panel H that is replaced by a locally
415 stable limit cycle in panel H'. These numerical simulations, for the parameter values used,
416 show that small and intermediate encounter rates make coexistence of both strategies in
417 polymorphic state less likely.

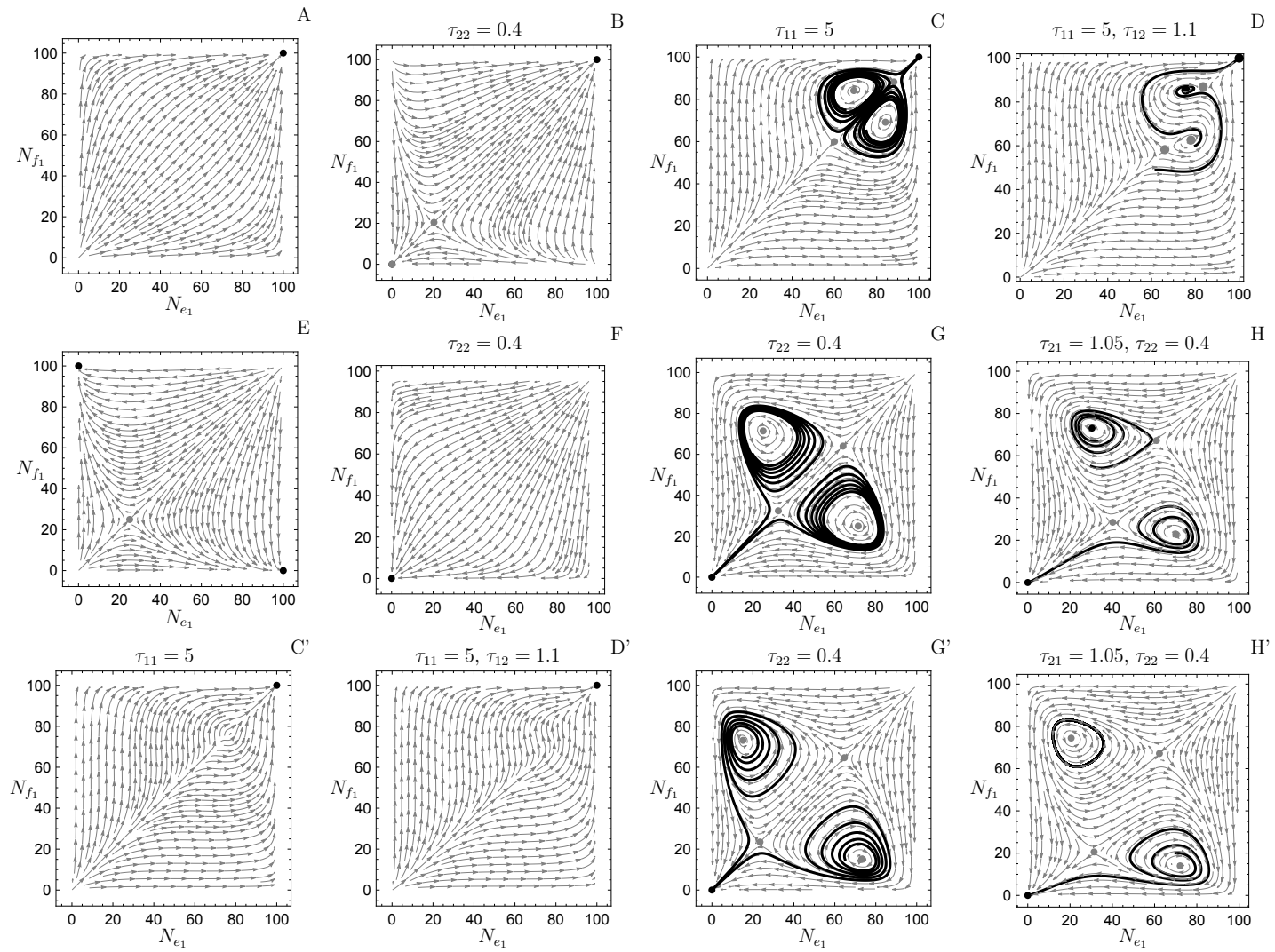


Figure 3: The replicator dynamics for the Owner–Intruder game when pairing is not instantaneous. For role-independent time-constrained bimatrix games (panels A, B, C, E, F, G, C', G'), the main diagonal remains invariant. The encounter rate between singles is $\lambda = 10$ in panels A–H and $\lambda = 1$ in panels C'–H'. Other parameters are the same as in the corresponding panels of Figure 2. Panels A and E are identical to their corresponding panels in Figure 2 since these are all equivalent to the classic bimatrix game. There are also no noticeable differences between panels B and F compared to Figure 2. The differences with Figure 2 (which emerges for very large λ) are as follows. For long interaction times between Hawks when $V > C$, the four interior NE of Figure 2 disappear completely when $\lambda = 1$ (panels C' and D') whereas the two asymmetric interior NE become unstable for intermediate λ (panel C). When the interaction time between Doves is short and $V < C$, the asymmetric interior NE of the role-independent time-constrained bimatrix game lose stability and the two symmetric interior shift apart as λ decreases (panels G and G'). With role-dependent interaction times, the asymptotically stable interior NE of Figure 2H eventually becomes unstable when λ decreases and a stable limit cycle emerges.

418 4. Discussion

419 This article extends to two-strategy bimatrix games the new approach to evolution-
 420 ary game theory developed by Křivan and Cressman (2017) for two-player, two-strategy,
 421 symmetric normal form games (i.e., matrix games) that incorporates the effect pair in-
 422 teraction times that depend on the players' strategies have on the evolutionary outcome.
 423 Evolutionary game theory applied to bimatrix games is based on two populations (or
 424 two roles) where individuals interact in pairs, one from each population. Classical bima-
 425 trix games, similarly to matrix games, assume that individuals get payoffs when paired,
 426 pairing is random and instantaneous, and the number of different types of pairs is given
 427 by the Hardy–Weinberg distribution. The evolutionary outcome of the bimatrix game is
 428 then predicted through an analysis of the NE structure of its payoff bimatrix and how
 429 this is connected to the eventual behavior of the game dynamics (e.g., the replicator
 430 equation). A complete analysis of the evolutionary outcome is well-known for all classical
 431 two-strategy bimatrix games (Hofbauer and Sigmund, 1998; Cressman, 2003).

432 When interaction times depend on strategies used by the pair, the Hardy–Weinberg
 433 distribution of pairs is no longer relevant and expected individual payoff is now a nonlinear
 434 function of the numbers using each strategy in the two populations whether the pair
 435 formation process among disbanded pairs is instantaneous (Section 2) or not (Section 3).
 436 However, in both cases, we show the existence of a unique distribution as a function of

437 these numbers at the beginning of these respective sections,¹² although we are only able
438 to provide an analytic expression for it when pair formation is instantaneous (see equation
439 (6)). Nevertheless, this allows us to define a time-constrained, bimatrix game in Section
440 2 and in Section 3 where payoff (which we call the fitness function) is given as expected
441 individual payoff per unit time. As pointed out in Sections 2 and 3.1, this new formulation
442 reduces to the classic bimatrix game when all interaction times are the same.

443 What is then of interest is how different interaction times affect the evolutionary
444 outcome. To this end, we completely characterized strict NE for all two-strategy, time-
445 constrained, bimatrix games (Sections 2.1 and 3.2 respectively). When pairing is in-
446 stantaneous (Sections 2.1) strict NE are characterized through their time-adjusted payoff
447 matrices (10). A strict NE corresponds to a locally asymptotically stable rest point of the
448 replicator equation where both populations use one of their pure strategies as indicated
449 by solid dots at vertices of the squares of Figures 2 and 3 respectively.

450 Unfortunately, other NE of the time-constrained bimatrix game are more difficult
451 to analyze. In particular, the analytic formula for interior NE is not available except
452 in special circumstances due to the complicated distribution that replaces the Hardy-
453 Weinberg distribution in these games. Since interior NE correspond to interior rest points
454 of the replicator equation, they can be approximated by simulating this dynamics for
455 particular games. No attempt is made for a complete analysis of all two-strategy time-
456 constrained bimatrix games.¹³ Instead, we focus on the time-constrained Owner-Intruder
457 game. This classic role-independent bimatrix game has an easily understood evolutionary
458 outcome.

459 When the cost of fighting over a resource C is less than its value V , both the owner of
460 the resource and the intruder should fight for it (i.e., both play Hawk) even though their
461 payoff by doing so is less than if they split the resource without fighting (i.e., both play
462 Dove) in the classic bimatrix game.¹⁴ The reason is that Hawk strictly dominates Dove
463 in each population. Although (Hawk, Hawk) remains a strict NE in the time-constrained
464 bimatrix game, other NE emerge as interaction times change. From panel B of Figures 2
465 and 3, we see that (Dove, Dove) can also be a strict NE (in which case there is also an
466 interior NE) when their interaction time is short enough compared to the equal time of

¹²In Section 3, this includes the distribution of pairs and singles

¹³The difficulty of doing such an analysis can be appreciated by considering the complete analysis for the two-locus two-allele viability selection model of population genetics. Pontz et al. (2018) show that this two-dimensional dynamics on the unit square has at least 192 different phase portraits. We feel our model will have a comparable (or even higher) number of different cases.

¹⁴The same result occurs for the bimatrix version of the Prisoner's Dilemma game where both players Defect at the evolutionary outcome even though they would be better off if both Cooperate.

467 the other interactions. Furthermore, while (Dove, Dove) is not a strict NE if only (Hawk,
468 Hawk) interaction time changes, up to four interior NE can appear if this interaction time
469 is large enough, some of which are (neutrally) stable and some unstable (panels C and
470 D).

471 When $V < C$, (Hawk, Hawk) is never a strict NE. In the classic bimatrix game,
472 there are two strict NE; namely, (Hawk, Dove) and the paradoxical ESS (Dove, Hawk)
473 where the intruder always wins the resource (i.e., the owner and intruder switch roles
474 through each interaction) as well as one unstable saddle symmetric interior NE where
475 both populations play the ESS of the classic symmetric Hawk–Dove matrix game. The
476 replicator equation predicts the paradoxical ESS will be the evolutionary outcome if and
477 only if the initial population distribution has more Hawks as intruders than as owners.
478 As shown in Figure 2, panels I and J, either one of these strict NE can disappear by
479 introducing a role dependence into the time interaction matrix (2). In fact, both must
480 disappear when (Dove, Dove) becomes a strict NE through their interaction time being
481 short enough compared to the equal time of the other interactions in Figures 2 and 3,
482 in which case interior NE may (panels G and H) or may not appear (panel F). There
483 are also marked differences between the evolutionary outcomes when pair formation is
484 instantaneous compared to when it is non instantaneous, as detailed in the main text.

485 In this article, although we have relaxed the implicit assumption of classic evolution-
486 ary game theory that all interactions take the same amount of time, we have assumed
487 that newly single individuals are immediately available to form pairs. This rules out
488 straightforward application of our methods to models where some single individuals from
489 a disbanded pair wait before joining the pair formation process. For instance, this occurs
490 in parental care models, e.g., Battle of the Sexes (Dawkins, 1976; Hofbauer and Sigmund,
491 1998; Mylius, 1999; Cressman, 2003; Broom and Rychtář, 2013) when males are immedi-
492 ately available to mate after a couple disbands whereas females will not mate immediately
493 but stay to care for offspring if abandoned by their mate. In this article, we assume that
494 both populations have the same number of individuals, which is required when pair for-
495 mation is instantaneous. On the other hand, when pair formation is non-instantaneous,
496 all calculations can be generalized to population 1 having a different size than population
497 2, although the formulas are more complex.¹⁵

498 In this article, we have generalised two-strategy bimatrix games by explicitly including
499 interaction times when pure strategists from each population are paired. When applied
500 to the classic Owner-Intruder game where each individual, at given interaction, is either a

¹⁵With unequal population size, the time-constrained bimatrix game with all τ_{ij} equal is no longer the classic bimatrix game as in Section 3.1.

501 Hawk or a Dove, we have a model where owners and intruders have a choice between two
502 levels of effort when engaged in a conflict (Hawks are willing to expend a great deal of time
503 and effort to obtain the resource while Doves are not). Another approach to this conflict
504 situation is to allow intermediate levels of effort, resulting in a time-constrained Owner-
505 Intruder game with a continuum of pure strategies. In the classic game with continuous
506 strategy sets (for a recent review see Cressman and Apaloo, 2018), the analysis of NE that
507 have additional properties such as Continuously Stable Strategy (CSS) or Neighborhood
508 Invader Strategy (NIS) are particularly important. Although beyond the scope of this
509 article, it is then essential to first understand the effect of interaction times on these
510 concepts of CSS and NIS.

511 The results of this article show that the evolutionary outcome for bimatrix games
512 becomes more complex when interaction times are incorporated into the game-theoretic
513 model. The results are also more complex than those reported by Křivan and Cressman
514 (2017) for matrix games with strategy-dependent interaction times as is to be expected
515 given the conceptual differences between classic matrix and bimatrix games. It is our
516 contention that these added complexities are often unavoidable to make the evolutionary
517 model more realistic. This is especially true when the model purports to describe a
518 behavioral system where pairs interact for different amounts of time.

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525 **Appendix A. Pairs distributional dynamics when pairing is instantaneous**

526 Here we derive pair dynamics (3). Let us consider a small time interval Δ . Because
 527 pairs n_{ij} split up following a Poisson process with parameter τ_{ij} , in this time interval
 528 a proportion $\frac{\Delta}{\tau_{ij}}$ of the n_{ij} pairs disbands and there will be $(\frac{n_{i1}}{\tau_{i1}} + \frac{n_{i2}}{\tau_{i2}})\Delta$ singles playing
 529 strategy e_i and $(\frac{n_{1j}}{\tau_{1j}} + \frac{n_{2j}}{\tau_{2j}})\Delta$ singles playing strategy f_j . The total number of disbanded
 530 singles in each population in time interval Δ is

$$\left(\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}} + \frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}} \right) \Delta. \quad (\text{A.1})$$

531 If these singles immediately and randomly pair, the proportion of newly formed n_{ij} pairs
 532 among all newly formed pairs will be

$$\frac{\left(\frac{n_{i1}}{\tau_{i1}} + \frac{n_{i2}}{\tau_{i2}} \right) \Delta}{\left(\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}} + \frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}} \right) \Delta} \frac{\left(\frac{n_{1j}}{\tau_{1j}} + \frac{n_{2j}}{\tau_{2j}} \right) \Delta}{\left(\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}} + \frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}} \right) \Delta}. \quad (\text{A.2})$$

533 To obtain the number of newly formed $(e_i f_j)$ pairs in the time interval Δ we multiply
 534 (A.2) by the number of newly formed pairs (which equals the number of disbanded singles
 535 because we assume instantaneous pairing) in time interval Δ and we obtain

$$\frac{\left(\frac{n_{i1}}{\tau_{i1}} + \frac{n_{i2}}{\tau_{i2}} \right) \left(\frac{n_{1j}}{\tau_{1j}} + \frac{n_{2j}}{\tau_{2j}} \right)}{\frac{n_{11}}{\tau_{11}} + \frac{n_{12}}{\tau_{12}} + \frac{n_{21}}{\tau_{21}} + \frac{n_{22}}{\tau_{22}}} \Delta.$$

536 Writing difference equations for pairs

$$n_{ij}(t + \Delta) = n_{ij}(t) - \frac{n_{ij}(t)}{\tau_{ij}} \Delta + \frac{\left(\frac{n_{i1}(t)}{\tau_{i1}} + \frac{n_{i2}(t)}{\tau_{i2}} \right) \left(\frac{n_{1j}(t)}{\tau_{1j}} + \frac{n_{2j}(t)}{\tau_{2j}} \right)}{\frac{n_{11}(t)}{\tau_{11}} + \frac{n_{12}(t)}{\tau_{12}} + \frac{n_{21}(t)}{\tau_{21}} + \frac{n_{22}(t)}{\tau_{22}}} \Delta \quad (\text{A.3})$$

537 and letting $\Delta \rightarrow 0_+$, we obtain the pair dynamics (3) in the main text.

538 From

$$\begin{aligned} N_{e_1} &= n_{11} + n_{12} \\ N_{f_1} &= n_{11} + n_{21} \\ N_{e_2} &= N - N_{e_1} \\ N_{f_2} &= N - N_{f_1} \end{aligned} \quad (\text{A.4})$$

539 and the generalized Hardy–Weinberg equation (5), Mathematica provides two equilibrium
 540 solutions for n_{ij} in terms of N , N_{e_1} and N_{f_1} . However, only the one given in (6) is non-
 541 negative when $0 \leq N_{e_1}, N_{f_1} \leq N$.

542 It is not immediately clear that $A \geq 0$ where A is given in (7). To see this, expand A
 543 as the following quadratic expression in N_{e_1}

$$A = N_{e_1}^2(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})^2 - 2N_{e_1}(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})(N\tau_{12}\tau_{21} - N_{f_1}(\tau_{11}\tau_{22} + \tau_{12}\tau_{21})) +$$

$$(\tau_{12}\tau_{21}(N - N_{f_1}) + N_{f_1}\tau_{11}\tau_{22})^2 = aN_{e_1}^2 + bN_{e_1} + c.$$

544 The minimum value of this upward parabola is

$$c - \frac{b^2}{2a} = 4N_{f_1}\tau_{11}\tau_{12}\tau_{21}\tau_{22}(N - N_{f_1}).$$

545 Since $0 \leq N_{f_1} \leq N$, this minimum is non-negative and so $A \geq 0$.

546 Appendix B. Calculation of the invasion fitness (16)

547 The fitness of strategy f_2 , Π_{f_2} , given in (9) calculated at the distributional equilibrium
 548 (6) is

$$\Pi_{f_2} = \frac{\sqrt{A}(\pi_{22}^f\tau_{12} - \pi_{12}^f\tau_{22}) + N\tau_{12}(\pi_{12}^f\tau_{21}\tau_{22} - 2\pi_{22}^f\tau_{11}\tau_{22} + \pi_{22}^f\tau_{12}\tau_{21})}{2\tau_{12}\tau_{22}(N - N_{f_1})(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})} -$$

$$\frac{(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})(\pi_{12}^f\tau_{22}(N_{f_1} - N_{e_1}) + \pi_{22}^f\tau_{12}(N_{e_1} + N_{f_1}))}{2\tau_{12}\tau_{22}(N - N_{f_1})(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})},$$

549 where A is given in (7). The invasion fitness of strategy f_2 when there are no individuals
 550 playing this strategy is then $\lim_{N_{f_1} \rightarrow N} \Pi_{f_2}$. We observe that

$$\lim_{N_{f_1} \rightarrow N} A = (N\tau_{11}\tau_{22} + N_{e_1}(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}))^2.$$

551 Since $N \geq N_{e_1}$, $N\tau_{11}\tau_{22} + N_{e_1}(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}) \geq 0$,

$$\lim_{N_{f_1} \rightarrow N} \sqrt{A} = N\tau_{11}\tau_{22} + N_{e_1}(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})$$

552 and the numerator of Π_{f_2} simplifies to

$$(N - N_{f_1})(\pi_{12}^f\tau_{22} + \pi_{22}^f\tau_{12})(\tau_{12}\tau_{21} - \tau_{11}\tau_{22}).$$

553 Thus, both the numerator and denominator of Π_{f_2} converge to 0 when $N_{f_1} \rightarrow N$ and we
 554 calculate the limit using L'Hospital's rule

$$\lim_{N_{f_1} \rightarrow N} \Pi_{f_2} = \frac{N_{e_1}\pi_{12}^f\tau_{21} + N_{e_2}\pi_{22}^f\tau_{11}}{N\tau_{11}\tau_{22} + N_{e_1}(\tau_{12}\tau_{21} - \tau_{11}\tau_{22})}. \quad (\text{B.1})$$

555 Similarly, the fitness of strategy e_2 , Π_{e_2} , given in (9) calculated at the distributional
 556 equilibrium (6) is

$$\Pi_{e_2} = \frac{\sqrt{A}(\pi_{22}^e \tau_{21} - \pi_{21}^e \tau_{22}) + N \tau_{21} (\pi_{21}^e \tau_{12} \tau_{22} - 2\pi_{22}^2 \tau_{11} \tau_{22} + \pi_{22}^2 \tau_{12} \tau_{21})}{2\tau_{21} \tau_{22} (N - N_{21}) (\tau_{12} \tau_{21} - \tau_{11} \tau_{22})} - \frac{(\tau_{12} \tau_{21} - \tau_{11} \tau_{22}) (\pi_{21}^e \tau_{22} (N_{e_1} - N_{f_1}) + \pi_{22}^e \tau_{21} (N_{e_1} + N_{f_1}))}{2\tau_{21} \tau_{22} (N - N_{e_1}) (\tau_{12} \tau_{21} - \tau_{11} \tau_{22})}.$$

557 The invasion fitness of strategy e_2 when there are no individuals playing this strategy is

$$\lim_{N_{e_1} \rightarrow N} \Pi_{e_2} = \frac{N_{f_1} \pi_{21}^e \tau_{12} + N_{f_2} \pi_{22}^e \tau_{11}}{N \tau_{11} \tau_{22} + N_{f_1} (\tau_{12} \tau_{21} - \tau_{11} \tau_{22})} \quad (\text{B.2})$$

558 by again applying L'Hospital's rule.

559 Appendix C. Uniqueness of distributional equilibrium of (25)

560 Fix N_{e_i} and N_{f_i} ($i = 1, 2$) and define $q_{e_i} = \frac{n_{e_i}}{N_{e_i}}$ ($q_{f_i} = \frac{n_{f_i}}{N_{f_i}}$) as the proportion of e_i (f_j)
 561 strategists in the population who are single. From (25) it follows that

$$\begin{aligned} q_{e_1} &= \frac{1}{1 + \lambda N_{f_1} q_{f_1} \tau_{11} + \lambda N_{f_2} q_{f_2} \tau_{12}} \\ q_{e_2} &= \frac{1}{1 + \lambda N_{f_1} q_{f_1} \tau_{21} + \lambda N_{f_2} q_{f_2} \tau_{22}} \\ q_{f_1} &= \frac{1}{1 + \lambda N_{e_1} q_{e_1} \tau_{11} + \lambda N_{e_2} q_{e_2} \tau_{21}} \\ q_{f_2} &= \frac{1}{1 + \lambda N_{e_1} q_{e_1} \tau_{12} + \lambda N_{e_2} q_{e_2} \tau_{22}}. \end{aligned} \quad (\text{C.1})$$

562 By Lemma 2 in Garay et al. (2017), there is a unique solution with q_{e_i} and q_{f_j} between 0
 563 and 1.

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