- **1** Physical constraints of cultural evolution of dialects in killer whales
- 2 Running title: Constraints of dialect evolution in killer whales
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Abstract Odontocete sounds are produced by two pairs of phonic lips situated in soft nares 24 25 below the blowhole; the right pair is larger and is more likely to produce clicks, while the left pair is more likely to produce whistles. This has important implications for the cultural 26 evolution of delphinid sounds: the greater the physical constraints, the greater is the 27 probability of random convergence. In this paper we examine the call structure of eight killer 28 whale populations to identify structural constraints and to determine if they are consistent 29 30 among all populations. Constraints were especially pronounced in two-voiced calls. In the calls of all eight populations, the lower component of two-voiced calls was typically centered 31 below 4 kHz, while the upper component was typically above that value. The lower 32 33 component of two-voiced calls had narrower frequency range than single-voiced calls in all populations. This may be because some single-voiced calls are homologous to the lower 34 component, while others are homologous to the higher component of two-voiced calls. 35 36 Physical constraints on call structure reduce the possible variation and increase the probability of random convergence, producing similar calls in different populations. 37

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40 I. INTRODUCTION

Understanding the physical basis of sound production is essential to categorize vocal signals 41 into natural categories. For the sounds of terrestrial mammals, the development of the source-42 43 filter theory has enabled researchers to describe the acoustic structure of sounds according to their mode of production and to predict acoustic variation caused by variation in the 44 anatomical or physiological attributes of the caller (Taylor and Reby, 2010). Even finer 45 resolution of phoneme classification in humans is based on the detailed knowledge of their 46 production: the position of tongue and lips, properties of the airflow and other features of the 47 48 vocal tract (Bickford, 2006).

49 Sounds of killer whales Orcinus orca have a complex structure (Ford, 1991; Yurk et al., 2002), but their categorization is hindered by the lack of understanding of their production 50 mechanisms. Odontocete sounds are not produced in the larynx, like in terrestrial mammals. 51 52 Instead, their source is situated in a complex system of air sacs and tissues that surround the nasal passage (Norris, 1968; Cranford et al., 1996; Cranford and Amundin, 2003). All toothed 53 whales (except sperm whales) have two pairs of phonic lips situated in each of the soft nares 54 55 on the ventral side of the vestibular air sacs, just below the blowhole; both pairs are used in sound production (Cranford et al., 1996; Cranford and Amundin, 2003). 56

Dolphin sounds were traditionally divided into three classes: clicks, burst-pulse sounds and 57 58 whistles. However, some species (e.g. killer whales, pilot whales *Globicephala sp.*, false 59 killer whales *Pseudorca crassidens*) regularly produce calls intermediate in structure between whistles and burst-pulse sounds, suggesting that these classes are not discrete, but rather the 60 two extremes of a perceptual continuum (Murray et al., 1998; Sayigh et al., 2013). Indeed, 61 Madsen et al. (2012) showed that dolphin 'whistles' are in fact not air-born, but produced by 62 pneumatically induced vibrations of phonic lips analogous to the operation of vocal folds in 63 terrestrial mammals. In most delphinoids the right pair of phonic lips is larger than the left 64

(Cranford et al., 1996). Madsen et al. (2013) examined the sound production of bottlenose
dolphin and false killer whale and showed that in both species clicks were produced by the
right pair while 'whistles' (tonal calls) were mostly produced by the left pair of the phonic lips.
This finding explains the long known observation that dolphins can simultaneously produce
clicks and 'whistles' (Murray et al., 1998).

However, some delphinoids can simultaneously produce two tonal sounds, yielding 'two-70 voiced' or 'biphonic' calls (e.g. killer whale: Ford, 1991; Tyson et al., 2007; pilot whale: 71 Savigh et al., 2013; bottlenose dolphin, *Tursiops truncatus*: Papale et al., 2015). Therefore, 72 the 'clicking' right pair of phonic lips is also capable of emitting tonal sounds. In addition, 73 74 Cranford et al. (2011) found that bottlenose dolphins can produce clicks with both pairs of 75 phonic lips working independently or simultaneously, though the right pair was used more often. So, the phonic lips are not strictly specialized, but differences in the size of the right 76 and left phonic lips suggest that they may be most effective for producing vocalizations with 77 different frequency characteristics. 78

This assumption has important implications for the cultural evolution of delphinid sounds. In 79 some, vocalizations are learned rather than transmitted genetically (killer whales: Ford, 1991; 80 81 Deecke et al., 2000; Foote et al., 2006; bottlenose dolphins: Tyack and Sayigh, 1997) and can indicate individuality (bottlenose dolphins: Janik and Savigh, 2013) or group affiliation (killer 82 whales: Ford, 1991). Killer whales have unique family dialects – sets of stereotyped sounds. 83 84 Calves learn their dialect from their mother and other family members (Bowles et al., 1988; Crance et al., 2014). With time, these dialects slowly change through learning errors and 85 innovations in a process called cultural evolution (Deecke et al., 2000; Wieland et al., 2010; 86 Filatova et al., 2015b). The recently diverged dialects are more similar than dialects that have 87 passed a long process of independent evolution (Ford, 1991; Deecke et al., 2010). However, 88 in some cases calls from distantly related dialects become more similar. Plausibly, this could 89

happen by either horizontal transmission or random convergence (Filatova et al., 2013). By 90 horizontal transmission we mean here the transmission of calls and call features from one 91 family to another. Random convergence occurs when calls of different families become more 92 93 similar by chance, without any influence on each other's evolution. This is more likely to happen if the variety of possible states is limited. In this case the variation in the diverging 94 95 calls soon reaches a limit and then the difference between them can no longer increase. This 96 situation is analogous to the phenomenon called "mutational saturation" which occurs when 97 many changes at a given locus eliminate phylogenetic signals (Delsuc et al., 2005). This happens because variation in nucleotide sequences is restricted by the structure of DNA: there 98 99 are only four nucleotides and four possible types of mutations (transitions, transversions, 100 insertions and deletions). Saturation can lead to homoplasy, i.e. convergence of traits in non-101 related taxa, which was described not only in genetics, but also in morphology (e.g. Alvarez et al., 1999; Mueller et al., 2004). 102

In vocal repertoires, a similar situation can be caused by constraints imposed on call structure by the physical properties of the sound producing apparatus. Therefore, it is important to understand these constraints to interpret the importance of call similarity: the stricter the constraints, the higher the probability that calls can become similar at random. In this paper we examine the call structure of eight killer whale populations to identify the scope of the constraints and test whether they are consistent among all populations.

109

110 **II. METHODS**

111 A. Data collection

Acoustic recordings used for this study were collected over various field projects using a
variety of equipment. All recordings were made at a sampling rate of 44.1 kHz or higher.

Recordings of the Northern resident and West Coast transient killer whale populations were 114 made from 1988 to 1999 in the waters off northern Vancouver Island. Recordings of the 115 Southern resident population were made from 1980 to 2009 in the waters of Salish Sea 116 117 (northern Washington State and southern Vancouver Island). The Alaskan resident killer whale population was recorded from 1984 to 2008 in Prince William Sound and Kenai Fjords 118 119 area. Calls from Kamchatkan residents were obtained in 2000-2014 off the southeastern coast 120 of Kamchatka peninsula. Eastern Aleutian transient killer whales were recorded in 2003-2008 121 in False Pass (between mainland Alaska and Unimak Island). Calls from Icelandic killer whales were obtained between 2008 and 2014 using both digital acoustic recording tags 122 123 (Dtags, Johnson and Tyack, 2003) and various other recording systems at two main locations: 124 one near Grundarfjörður on the Snæfellsnes Peninsula (West Iceland) the other near Vestmannaeyjar off the southern coast of Iceland. Calls from Norwegian killer whales were 125 obtained between 2005-2009 using Dtags in the northern Norwegian Vestfjord fjord system 126 and off Vesterållen. 127

Calls from four North Pacific resident populations and the West Coast transient population
were classified according to existing catalogues (Ford, 1987; Yurk et al., 2002; Filatova et al.,
2004). For the False Pass transient population, call categorisation was performed based on
audible characteristics of calls and visual inspection of spectrographic features. For Icelandic
and Norwegian killer whales, we have modified and updated existing catalogues (Strager,
1995; Shapiro, 2008; Duc, 2010).

Each killer whale population produces several tens of stereotyped call types recognized even
by inexperienced observers (Yurk et al., 2002; Shamir et al., 2014); these call types retain
stereotypy in different circumstances (Ford, 1989) and over the years (Deecke et al., 2000;
Foote et al., 2008). Call types have different degrees of structural variation: some types are
very stereotyped while others demonstrate some variation that can be either discrete or

gradual. Call types with a discrete variation are traditionally divided into a number of 139 subtypes according to their structural characteristics; call types with a gradual variation are 140 141 not divided into subtypes, and all variations are regarded as a single call type (Ford, 1991). To 142 cover the full range of structural variation within a type, we selected two calls from each call 143 type or subtype that were the least similar to each other. If a call type had no subtypes, two 144 samples from that call type were used; for call types that comprised discrete subtypes, we 145 used two samples from each subtype. For calls that did not fall into discrete subtypes but 146 showed apparent variations, two calls from opposite extremes of the structural continuum 147 were selected. When possible, the pairs of call samples from the same type/subtype were 148 selected from different encounters and different years to cover the presumed variation in the call structure. 149

Norwegian killer whales are known to produce compound calls, which consist of
combinations of other stereotyped calls produced in stable sequences (Strager, 1995; Shapiro
et al., 2011). Treating every stable sequence as a separate type would produce pseudoreplication through the increased presence of calls used as parts of stable sequences (as they
will be measured twice – as a separate call and as a part of a stable sequence). Thus, we only
measured those stable sequences that contained calls or syllables not observed occurring
separately.

In total, 638 call samples were used for the analysis: 96 from Alaskan residents, 102 calls
from Kamchatkan residents, 96 calls from Northern residents, 62 calls from Southern
residents, 34 calls from Eastern Aleutian transients, 36 calls from West Coast transients, 72
calls from Norway, and 140 calls from Iceland. Sample sizes of calls from the transient
populations were lower because the overall repertoire size of stereotyped calls is typically
lower in these populations (Ford, 1987; Saulitis et al., 2005). The differences in sample sizes

163 from different populations did not bias our results, because each sample size was selected to 164 cover the full range of structural variation of stereotyped calls in the respective population.

165 **B. Acoustic and statistical analysis**

To capture the frequency parameters over the whole call duration, we extracted call contours following the method described in Filatova et al. (2012). The extracted contours represented a set of frequency measurements of each call's fundamental frequency spaced 0.01 s apart. If fundamental frequency was not visible, we measured one of the harmonics and divided the measurements by the number of that harmonic to obtain the fundamental frequency values. For two-voiced (biphonic) calls, containing overlapping frequency components, we extracted the contours of both the lower- and upper-frequency components.

173 In many two-voiced calls the start and end of the lower and upper components do not match,

so that a call can be partly two-voiced and partly single-voiced (Fig. 1). Single-voiced

segments can consist of either lower or upper component (Fig. 1). As we were interested in

176 constraints imposed by the structure of both pairs of phonic lips, we analysed two-voiced and

single-voiced segments of the calls separately. For each point of a two-voiced segment of

each two-voiced call we obtained two values: frequency of the lower and the upper

179 components at that point. Single-voiced segments of two-voiced calls (both from the lower

180 and upper components) were pooled together with single-voiced calls; for each point of these

181 vocalizations we obtained a single frequency measurement.

182 We analyzed the distribution of frequency values of the lower and the upper components of

two-voiced call segments, and in single-voiced calls and call segments using plots and

184 descriptive statistics in R (R Core Team, 2014).

185

186 **III. RESULTS**

To identify the scope of the constraints of the sound producing apparatus in killer whales, we 187 analyzed the frequency range of two-voiced and single-voiced calls and call segments in 188 189 different populations. In the two-voiced segments of calls the frequency of the lower 190 component was typically below 4 kHz (Fig. 2). Only in five calls the frequency values of the lower component ranged above 4 kHz. The first call type with frequency of the lower 191 192 component ranging above 4 kHz was AKS16b from the Alaskan resident population. The 193 lower component of this call is a short upsweep that sometimes can rise above 4 kHz in the 194 end. In our sample, only one frequency point of one AKS16b call ranged above this value; the 195 second call of this type had the whole lower component below 4 kHz.

Four other calls with the frequency values of the lower component above 4 kHz belonged to two call types: the AKS18 call type from the Alaskan resident population and the I84 call type from the Icelandic population (Fig. 3). The most of the lower component of both AKS18 calls and about the half of the contour of the lower component of both I84 calls were centered above 4 kHz.

The frequency of the upper component was typically centered above 4 kHz, though in some 201 202 calls the upper component ranged down to 3 kHz (Fig. 2). The upper component ranging 203 below 4 kHz was found in sixteen call types from all populations except Norwegian. Usually (in eleven of the sixteen call types) it occurred in calls where the upper component started 204 205 with an abrupt upsweep, and its section with frequency below 4 kHz was rather short – 10-30 206 ms. Only five call types included relatively long (> 50 ms) sections of the upper component with frequency below 4 kHz: K20 call type from Kamchatkan residents, N18 call type from 207 208 Northern residents, T12ii call type from West Coast transients, FP4 call type from Eastern 209 Aleutian transients, and I41 call type from Iceland.

The frequency of the lower component of two-voiced call segments had narrower range thansingle-voiced calls and call segments in all populations (Fig. 4). The 5% quantile of single-

212	voiced calls was lower than the 5% quantile of the lower component of two-voiced call
213	segments in all populations except West Coast transients, and the 95% quantile of single-
214	voiced calls was higher than the 95% quantile of the lower component of two-voiced call
215	segments in all populations, except Alaska (Table 1).
216	The 95% quantile of the lower component was very similar among all populations except
210	The 35% quantite of the lower component was very similar among an populations except
217	transients (that had lower values) and Alaskan residents that had a much higher value due to
218	the presence of a single call type (AKS18). The 5% quantile of the lower component was
219	more variable, but in all populations except West Coast transients it was above 300 Hz.
220	All populations had similar 5% and 95% quantiles for the upper frequency component, except
221	for West Coast and Eastern Aleutian transients that had lower values. Nevertheless, in all
222	populations the 95% quantile for single-voiced calls was lower than the 95% quantile of the
223	upper component of two-voiced call segments (Table 1).

224

225 IV. DISCUSSION

In the stereotyped calls of all eight killer whale populations, the frequency values of the lower 226 227 component were mostly centered below 4 kHz, while the frequency values of the upper 228 component were mostly above that value. It appears that 4 kHz is a natural boundary between 229 lower and upper components. Killer whales are technically capable of producing lower 230 components at frequencies above, and upper components at frequencies below this value, but 231 such calls are rare, suggesting that there is a physical constraint. Despite the relatively small 232 sample size (two calls per type/subtype, total of 638 calls from eight populations), the overall 233 frequency values of two-voiced calls from different populations were mostly similar (Table 1), suggesting that the sample size was enough to cover the full range of variations in 234 235 stereotyped calls of the studied populations.

236

A. Implications for sound production

237 Many odontocete species have been reported to produce two overlapping sounds simultaneously (e.g., bottlenose dolphin: Lilly and Miller, 1961; Risso's dolphin: Corkeron 238 239 and Van Parijs, 2001; common dolphin: Moore and Ridgway, 1995; short-finned pilot whale: 240 Sayigh et al., 2013; long-finned pilot whale: Nemiroff and Whitehead, 2009; false killer whale: Murray et al., 1998; beluga whale: Belikov and Bel'kovich, 2006; Garland et al., 241 242 2015), but the structure of these sounds varies across species. An upper component is typically a high-frequency tonal sound ('whistle') in all species, but most dolphins (bottlenose 243 244 dolphin, common dolphin, Risso's dolphin) and beluga whales typically produce burst-pulse 245 sounds or click trains as a lower component, and only killer whales, false killer whales and 246 both species of pilot whales have been reported to regularly produce calls as a lower component of two-voiced sounds. 247

Research on sound production in odontocetes has been so far mostly focused on bottlenose 248 249 dolphins that typically produce click trains (echolocation clicks and burst pulse sounds) and high-frequency whistles, but rarely emit "calls" - the sound category intermediate between 250 251 click trains and whistles (Murray et al., 1998). Even in the study of sound production where a 252 false killer whale Pseudorca crassidens was involved (Madsen et al., 2013), the authors 253 referred to the tonal vocalizations as "whistles" and did not provide any sonograms that could 254 clarify the structure of these sounds. In many cases, the terminology used to classify sounds 255 into categories is still poorly linked to sound production mechanisms, often due to a lack of 256 knowledge on the mechanisms involved in producing different types of sound. In the future, advances in the understanding of delphinid sound production would benefit from 257 258 collaboration between morphologists and bioacousticians familiar with vocal repertoires. 259 The frequency of the lower component of two-voiced call segments had narrower range than

single-voiced calls and call segments in all populations: single-voiced calls had lower

261 minimum and higher maximum frequency values. The higher maximum frequency values are 262 easy to explain: we did not make a distinction between upper and lower components in single-263 voiced calls and call segments, so some single-voiced calls and call parts are homologous to 264 the lower component while others are homologous to the upper component of two-voiced 265 calls. This distinction is important to consider in future studies of killer whale acoustic 266 behavior, because "upper" and "lower" single-voiced calls can potentially have different 267 communicative functions.

It is less clear why single-voiced calls had lower minimum values than the lower component 268 of two-voiced calls. In our dataset, the 5% quantile of the lower component in two-voiced 269 270 calls was above 300 Hz in all populations except West Coast transients. In West Coast 271 transients, the lower values of the lower component were due to the presence of WCT12 call type that has a two-voiced segment with the frequency of the lower component below 300 Hz 272 (Fig. 5). However, the non-harmonic frequency modulation in this segment suggests that the 273 sidebands are not true harmonics. In humans, a similar phenomenon occurs when both vocal 274 folds vibrate at different frequencies (see Fig. 3 in Wilden et al., 1998). 275

276 Therefore, our results indicate that killer whales rarely produce two-voiced calls with a lower 277 component below 300 Hz, though they often produce sounds with frequency below 300 Hz separately as single-voiced calls. One of the reasons can be that it is physically difficult to 278 produce two sounds with so drastically different structure simultaneously. However, dolphins 279 280 and beluga whales routinely produce simultaneous whistles and burst pulse sounds (Lilly and Miller, 1961; Corkeron and Van Parijs, 2001; Moore and Ridgway, 1995; Van Parijs, 2001; 281 282 Garland et al., 2015). Dolphin burst pulse sounds resemble low-frequency killer whale calls and likely result from a similar sound production mechanism (Murray et al., 1998). 283

Dolphin sounds are produced by two pairs of phonic lips (Cranford et al., 2011; Madsen et al.,

285 2013). Two-voiced calls most likely arise from both pairs of phonic lips oscillating

simultaneously with different frequencies, one pair producing the upper and another – the
lower component. In this case, the lack of the upper component in calls below 300 Hz can
indicate that both pairs of phonic lips are involved in their production (i.e., technically the
low-frequency sounds are two-voiced, but both pairs of lips oscillate with low frequency).

It is also possible that sound sources other than phonic lips might exist. Most two-voiced 290 calls of killer whales have heterodyne frequencies below and above the upper component 291 292 (Fig. 1). Heterodyne frequencies arise from the interaction of the lower and upper components 293 (Wilden et al., 1998; Brown, 2008), confirming that these components are produced by coupled sound sources (most likely, two pairs of phonic lips). However, some call types lack 294 295 the heterodynes, suggesting either that the level of coupling between the right and the left 296 phonic lips pairs can vary across call types, or that an alternative sound production source can be involved. Production of two-voiced calls by baleen whales (Gedamke et al., 2001; Tyson et 297 298 al., 2007; Tervo et al., 2011) that lack the system of phonic lips also suggests that cetaceans 299 can have other potential sources of sound production.

Terrestrial mammals can produce biphonic calls without specific anatomical adaptations of 300 301 the sound-producing structures (Frey et al., 2016). Humans produce sounds with two 302 independent, but similar frequencies by asynchronous vibration of the left and right vocal folds (Tigges et al., 1997). Other proposed sources of biphonation in terrestrial mammals 303 304 include air vortices at the narrowings of the vocal tract (Solomon et al., 1995) and source-305 filter interaction when the vocal folds start oscillating at one of the resonance frequencies of the vocal tract (Titze et al., 2008; Volodin et al., 2013). Both of these mechanisms are 306 307 however unlikely to occur in cetaceans because these mechanisms relate on air resonances depending on the air volume in the vocal tract. This would lead to the shift in the fundamental 308 309 frequency when the whales dive and air cavities contract under pressure. However, both lower 310 and upper components of stereotyped two-voiced calls in killer whales are produced with

stable fundamental frequencies (Ford, 1991; Yurk et al., 2002; Miller et al., 2007). Miller et
al. (2007) demonstrated the differences in the relative intensity of harmonics between male
and female killer whales, suggesting that air volume in odontocete's nasal sacs can act as a
filter similarly to the vocal tract in terrestrial mammals.

If a sound source besides right and left pairs of phonic lips exists, three-voiced sounds are 315 316 theoretically possible. Such sounds have not been reported in killer whales, but Sayigh et al. 317 (2013) provides a sonogram (Fig. 2C and Fig. 5 in Sayigh et al., 2013) of a call of short-318 finned pilot whale *Globicephala macrorhynchus* that apparently contains three independently modulated components: an upsweep upper component, a downsweep lower component and a 319 320 low-frequency buzz or squeak. The upper component has clear heterodynes, indicating that 321 the upper and lower components are produced by coupled sources, most likely phonic lips, but the source for the overlapping low-frequency buzz is unknown. Future studies are 322 required to understand the sound production of two-voiced (and possibly three-voiced) calls 323 in killer and pilot whales using similar experimental approaches as used in bottlenose 324 dolphins. 325

326 We found that constraints were especially pronounced in two-voiced calls. Single-voiced calls 327 ranged further down than the lower component of two-voiced calls. Upper component ranged higher than single-voiced calls; however, in this study we have not considered so-called 328 'whistles' that can occur at much higher frequencies (Samarra et al., 2010; Filatova et al., 329 330 2012b; Simonis et al., 2012). Whistles were not a part of this study because we included only stereotyped calls that allow consistent sampling, while whistles are typically not stereotyped 331 332 (Ford, 1991). Stereotyped whistles were reported in killer whales, but they were less common than calls and not specific to a particular pod or clan (Riesch et al., 2006). This suggests that 333 334 they have other communicative function than stereotyped calls, probably due to the 335 differences in their propagation range (Thomsen et al., 2002; Miller, 2006).

Besides stereotyped calls and whistles, killer whale acoustic repertoire includes variable calls that cannot be divided into discrete categories and so-called aberrant calls that represent highly distorted stereotyped calls. Analyzing the structure of these calls as well as whistles can provide important insights into sound production mechanisms in killer whales.

340

B. Implications for cultural evolution

Calls with either lower or upper component frequencies that go far beyond the assumed 341 boundary of 4 kHz were rare in general and completely absent in some populations. Calls 342 with the frequencies of the lower component above 4 kHz were found only in the Alaskan and 343 344 Icelandic populations. Of note, Alaskan AKS18 call has sidebands (Fig. 3) in some (but not 345 all) samples, suggesting that the real fundamental frequency can be lower, and the energy is shifted to the upper harmonic due to filtering. The lower limit of the upper component is not 346 347 as strict, but only in transient populations the 5% quantile of the upper component goes below 4 kHz, confirming that transients generally have lower call frequencies than residents and 348 349 North Atlantic killer whales (Filatova et al., 2015a). Besides, the lower component in twovoiced calls rarely goes below 300 Hz, though killer whales often produce such low-350 351 frequency sounds separately as single-voiced calls.

If some socially learned sounds are easier to produce than others, the morphology of the 352 353 phonic lips plays a role in the formation of a cultural attractor. The idea of cultural attraction 354 (Sperber, 1996) is based on two contradicting observations: a) social learning is generally not a copying process and typically results in modifications of the transmitted information or 355 behavior; but b) cultural information/behavior is often relatively stable within whole 356 populations and across generations. Cultural attractors are abstract 'centers of gravity' in the 357 space of possibilities, which exist because some factors affect the probability that individual 358 359 memes will depart from their models in one direction rather than in another. In killer whales, it appears that such 'centers of gravity' occur at frequencies around 1 kHz for the lower 360

361 component and around 5-10 kHz for the upper component of two-voiced calls. This would
362 define the most probable direction of call change, causing multiple convergences in different
363 populations.

This finding is particularly important for the understanding of the cultural evolution of killer whale sounds. Physical constraints on call structure reduce the possible variation, so that calls more often become similar by random convergence. Indeed, very similar calls were found in different populations (Fig. 6). Since these populations are not in contact, and some of them are separated by thousands of kilometers, the most plausible explanation is random convergence.

370 Filatova et al. (2012a) found no correlation between dynamic time warping similarity of calls and geographic distance in four North Pacific resident populations. One of the suggested 371 372 explanations of this finding was that calls change too fast and populations soon reach the 373 maximum possible divergence, and after that they can only converge, so the similarity of 374 population repertoires is not phylogenetically meaningful (at least, at the level of dynamic time warping comparison of call contours). The current study provides clear evidence in favor 375 376 of this hypothesis. A similar situation exists in human languages: phonemes change faster 377 than other language features (Labov, 2011), but due to the limited variation they often 378 converge randomly in non-related languages (Moran et al., 2014).

This phenomenon is especially important to understand while interpreting the similarities in calls of captive-born killer whales. For example, Kremers et al. (2012) found some vague similarities of calls of captive-born killer whales of Icelandic maternal origin with calls of Canadian Northern and Southern residents (even though none of them was related to Northern residents, and only two of the four whales had Southern resident grandfathers). Kremers et al. (2012) explain this finding by multiple chain learning events among captive individuals, ignoring the arguably more plausible explanation of random call convergence.

Sounds of killer whales and other cetaceans are usually referred to as culturally transmitted, as opposed to genetically inherited sounds of most other mammals. However, Laland and Janik (2006) emphasize that it is counterproductive to interpret behavior as being either genetic or cultural, because every learned behavior has some genetic basis. This basis can include the tendency to learn specific behaviors as well as physical ability to perform them. Our work illustrates the genetically inherited constraints of socially learned killer whale sounds that shall be considered in the further studies of their cultural evolution.

393

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Table 1. 5% and 95% quantiles of frequency values (in kHz) of single-voiced and both

	Single-voiced		LFC of two- voiced		UFC of two- voiced	
	5%	95%	5%	95%	5%	95%
Kamchatka	0.13	6.55	0.42	1.92	5.37	10.76
Alaska	0.10	5.63	0.34	5.71	5.04	10.64
Northern residents	0.05	4.63	0.54	2.17	5.12	10.66
Southern residents	0.04	4.80	0.36	2.02	4.51	8.12
West Coast transients	0.23	2.99	0.20	0.90	3.32	6.71
Eastern Aleutian transients	0.25	4.73	0.40	1.49	3.28	5.42
Iceland	0.25	4.33	0.66	2.14	5.01	10.47
Norway	0.27	7.52	0.32	2.50	5.21	11.28

577 components of two-voiced calls and call segments in eight studied populations.

578

580 Figure captions

581 Fig. 1. (color online) The start and the end of the lower-frequency component (LFC) and

upper-frequency component (UFC) in two-voiced call do not match, so that only the middle

segment of the call is two-voiced, while the beginning and the end of the call are single-

- voiced. Heterodyne frequencies arise from the interaction of LFC and UFC; the lower
- heterodyne in each point is equal to LFC-UFC, and the higher heterodyne is equal to

586 LFC+HFC.

- 587 Fig. 2. (color online) Scatterplot with marginal histograms showing the distribution of
- 588 frequency points of the lower and upper components in two-voiced calls from all populations:
- 589 KR Kamchatkan residents, AR Alaskan residents, NR Northern residents, SR –
- 590 Southern residents, WT West Coast transients, AT Eastern Aleutian transients, Ice –
- 591 Iceland, Nrw Norway.
- Fig. 3. Two call types with the lower component above 4 kHz: Alaskan call AKS18 (left) andIcelandic call I84 (right).
- Fig. 4. (color online) Density plots of the frequency values of the lower (blue) and upper
- 595 (green) components of two-voiced call segments and of single-voiced calls and call segments
- 596 (red) from all studied populations.
- Fig. 5. (color online) WCT12 call type showing the two-voiced segment with the frequency of
 the lower component below 300 Hz. Note the non-harmonic modulation in the beginning and
 the end of the segment (marked by arrows) indicating that the sidebands are not true
 harmonics.
- Fig. 6. Examples of convergence of call structure in different populations.

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