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Local thermal non-equilibrium in sediments: Implications for temperature dynamics and the use of heat as a tracer

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34 Abstract

35 Understanding streambed thermal processes is of fundamental importance due to the effects of temperature dynamics on stream ecology and solute exchange processes. Local Thermal 36 37 Equilibrium (LTE) between fluid and solid is usually assumed for modelling heat exchange in streambeds and for inferring pore water flow velocities from streambed temperature data. By 38 examining well established experimental and theoretical relationships of the fluid-solid heat 39 40 transfer coefficient in a numerical scheme for a range of Reynolds (Re) numbers (0.01 > Re >41 0.001), we show here that, for a range of typical streambed conditions, LTE is not attained. Thus errors in velocity estimates obtained when inverting streambed temperature data assuming LTE 42 can be considerable especially at relatively low flow rates. We show that for certain conditions 43 were the LTE assumption is not valid, inferred pore water velocities of up to 1 m/d can be 44 obtained with LTE assumption even if the actual velocities are much smaller or even zero. 45 46 Ignoring the possibility of Local Thermal Non-Equilibrium (LTNE) will have consequences for the correct estimation of streambed pore water and heat fluxes at low Re values. More 47 laboratory studies are urgently needed to supplement the sparse existing data in this area and 48 further test the findings of this study. 49

50 Keywords: Local thermal non-equilibrium, Heat as a tracer, Heat transfer, streambed

51

52 **1. Introduction**

Understanding streambed temperature dynamics is critical to deriving deeper insights into 53 stream ecology. Temperature is a fundamental biological variable and is a major control on 54 biogeochemical processes which underpin vital ecosystem services [1]. 55 Moreover. measurements of temperature variability between streams and groundwater [2] can be used to 56 57 infer patterns and processes of hyporheic exchange [3] and are critical for controlling nutrient and carbon cycling in streambed systems and the potential attenuation of contaminants in the 58 hyporheic zone [4]. Most techniques which use heat as a tracer rely on a physically based 59 model which inverts temperature measurements to infer flow rates and sediment thermal 60 61 properties [5]. The most popular methods take advantage of the solar signal which generally 62 induces heat exchange between streams and underlying sediments [6-9]. A damping and 63 attenuation of the diel stream temperature signal with depth is normally observed and most 64 methods assume a 1-D flow field for interpretation, although recent studies have shown that this

may be problematic in real, non-uniform, flow fields [10, 11]. Additional uncertainties may
stem from sediment heterogeneity [12], measurement error and difficulties in estimating thermal
parameters [13, 14].

Despite its increasing popularity in the hydrological community, all studies to date which have 68 used heat as a tracer for investigating groundwater-surface water interactions in streambed 69 environments have assumed the validity of the single-temperature (i.e. using a single domain to 70 71 model temperatures for the solid and fluid in combination) heat transport equation. This relies 72 on the assumption of instantaneous local thermal equilibrium between the solid matrix materials 73 and the pore fluids. However, we show here, by drawing on the extensive literature on this subject from other fields and proposing a new correlation, that this assumption is questionable in 74 75 the context of many streambeds. As a result, considerable errors in flux estimation and 76 conceptual understanding of streambed thermal processes may result.

77

78 **2. Methods**

79 2.1 Deriving the heat transfer coefficient at low Reynolds numbers typical of streambeds

When the assumption of LTE is suspected to break down, the temperature of solid and fluid 80 phases have to be considered separately rather than as a single average temperature field. In this 81 82 two-domain approach, it is assumed that each phase is continuous and represented by an appropriate effective total thermal conductivity and therefore effective thermal diffusivity [15, 83 16]. We use a Dispersion-Particle-Based two-equation model based on the heat transfer 84 coefficient between the solid and the fluid phases. The equations for the solid and the fluid 85 phases without heat sources or sinks and without an energy term for viscous-work can be 86 expressed as [17, 18]: 87

(1)

(2)

88
$$\frac{\partial T^{f}}{\partial t} + v \frac{\partial T^{f}}{\partial x} = \frac{1}{\phi} \left(\frac{k_{f}}{(\rho c_{p})_{f}} + f\left(\vec{\beta}, \vec{v}\right) \right) \frac{\partial^{2} T^{f}}{\partial x^{2}} + \frac{h_{sf} a_{sf}}{\phi(\rho c_{p})_{f}} (T^{s} - T^{f})$$

89
$$\frac{\partial T^s}{\partial t} = \frac{k_s}{(1-\phi)(\rho c_p)_s} \frac{\partial^2 T^s}{\partial x^2} - \frac{h_{sf}a_{sf}}{(1-\phi)(\rho c_p)_s} (T^s - T^f)$$

Where, a_{sf} is the surface area of particle per unit volume of porous media, h_{sf} is the heat transfer coefficient, ϕ is the overall porosity and k is the thermal conductivity tensor, respectively where f represents the fluid phase and s represent the solid phase. Also $(\rho c_p)_f$ is

93

the volumetric heat capacity of fluid, $(\rho c_p)_s$ is the volumetric heat capacity of solid, T^f is the

fluid temperature, T^s is the solid temperature and t is the time. In addition $f(\vec{\beta}, \vec{v})$ is the 94

95 hydrodynamic dispersion function:
$$f(\vec{\vec{\beta}}, \vec{v}) = \vec{\vec{\beta}} \cdot \left(\frac{\rho_f c_f}{\rho c} \cdot \vec{v}\right)^2$$
 proposed by Rau et al. [19]

where $\vec{\beta}$ is the thermal dispersivity matrix and \vec{v} is the average pore water velocity defined as a 96 vector. In this form of the hydrodynamic dispersion function the thermal dispersivity has the 97 units of [T]. In Eqs. 1 and 2, the surface area of particles per unit volume of porous media can be 98 99 estimated by [20]:

100
$$a_{sf} = \frac{6(1-\phi)}{dp} \tag{3}$$

Where, dp is defined as the average grain size of the porous media as would be obtained from a 101 grain size distribution curve. It should be noted that this equation may not be valid for poorly 102 sorted sediment, but is applicable to the homogeneous conditions modelled here. In order to 103 determine the heat transfer coefficient between the fluid and solid particles, a number of 104 experimental correlations have been proposed [21-23]. However, despite extensive effort, no 105 theory has been developed which can satisfactorily describe the heat transfer rate over a wide 106 range of porous media with different physical properties, such as grain size or velocity 107 distribution [21]. At high Reynolds numbers, there is a well-accepted correlation which has 108 been used to solve the heat transfer in porous beds for more than three decades. It is expressed 109 110 as [21]:

111
$$Nu = 2 + 1.1 \operatorname{Pr}^{\overline{3}} \operatorname{Re}^{0.6}$$
 (4)

where, Nu, Pr and Re are the dimensionless Nusselt number, Prandtl number and Reynolds 112 number defined as: 113

114
$$Nu = \frac{h_{sf}dp}{k_f}, \text{ Pr} = \frac{c_{pf}\mu_f}{k_f}, \text{ Re} = \frac{\rho_f v dp}{\mu_f}$$
 (5)

where, μ_f , $c_{_{pf}}$ and ρ_f are the fluid viscosity, fluid heat capacity and fluid density. Increase in 115 Re enhances heat and momentum transfer between fluid particles which increases the friction 116

force on the grain surface and therefore the heat transfer rate. The average grain thermal Peclet number (Pe_{ave}) describes the ratio of the advective to conductive heat transport and defined as:

119
$$Pe_{avg} = \frac{\rho_f c_{pf} v dp}{k_e}$$
(6)

where, k_e is the average heat conductivity of the porous medium defined as $k_e = k_s^{(1-\phi)} k_f^{(\phi)}$. The 120 proposed correlation (equation 4) explains the experimental data obtained by many authors [24, 121 122 25] for Re>1. However, such high Re are not expected in streambeds unless the grainsize and thus hydraulic conductivity of the bed are sufficiently great and large hydraulic gradients are 123 also present to drive high fluid velocities such as might be the case in high energy losing stream 124 systems [26]. For example, a gravel streambed with an average grainsize of 1 mm and a pore-125 water velocity of 10 m/d would have a Re of around 0.1 ($Pe_{ave}=0.074$ when $k_s=2.5$ W(mC)⁻¹). 126 127 However, many streambed environments have smaller grain sizes (silt to sand i.e. 0.01 mm to 1 mm) or smaller pore water velocities due to lower ambient hydraulic gradients such as are 128 often found in lowland settings [7, 14, 27] leading to relatively low Reynolds numbers. For 129 example a sandy streambed (dp=0.3 mm) with a pore water velocity of around 0.3 m/d would 130 have a *Re* of approximately 0.001 ($Pe_{avg}=7.4 \times 10^{-4}$ when $k_s=2.5$ W(mC)⁻¹). 131

For *Re*<1 relevant to many streambed environments, fewer data are available and equation (4) 132 133 breaks down. Therefore, we propose a correlation based on the only experimental data published to date [28] to calculate the heat transfer coefficient at low Reynolds numbers (down 134 to Re=0.001). These data have been widely used in various studies in the literature [22, 29, 30]. 135 In order to obtain a correlation of the heat transfer in saturated sand, only the part of the Kunii 136 and Smith [28] data related to experimentation with water as the fluid phase and sand and glass 137 beads (with thermal conductivity of 0.5 $W(mC)^{-1}$) as the solid phase were plotted and analysed 138 (Figure 1). The mathematical equation explaining the physics of heat transfer of a single sphere 139 submerged in a fluid is used as the basis of the analysis [31]: 140

141
$$Nu = 2.0 + K_1 \operatorname{Pr}^p \operatorname{Re}^q$$

where, K_1 , p and q are experimental coefficients. It is discussed in Nelson and Galloway [22] that the coefficient of 2 in equation (7) is only valid for single sphere and this coefficient needs to be measured experimentally for real materials. It is also shown by Lienhard [32] that the ratio of thickness of the thermal boundary layer δ_t to that of the fluid boundary layer δ_f equals to:

(7)

146 $\frac{\delta_t}{\delta_f} = \Pr^{\frac{-1}{3}}$ for a wide range of gas and fluids $0.6 \le \Pr \le 50$. Thus, in derivation of the heat

transfer equation the Prandtl number takes the power of 1/3. Therefore, we would expectequation (7) to take the following form:

(8)

(9)

149
$$Nu = \alpha + K_1 \operatorname{Pr}^{\overline{3}} \operatorname{Re}$$

We used the software *Datafit* to fit equation (8) to the Kunii and Smith [28] experimental data by varying the parameters α , K_1 and q by a least squares method. The coefficients were chosen from the best fit (details of fitting parameters and confidence intervals can be found in Table A and B in Appendix A). In addition, the model proposed by Nelson and Galloway [22] is also considered to compare the results of each model at *Re*=0.01. The Nelson and Galloway model has been widely used in the industry applications having Reynolds numbers down to 0.01 [33, 34]. The model has the form:

157
$$Nu = \frac{2\varsigma + \left(\frac{2\varsigma^2(1-\phi)^{1/3}}{[1-(1-\phi)^{1/3}]^2} - 2\right) \tanh \varsigma}{\frac{\varsigma}{1-(1-\phi)^{1/3}} - \tanh \varsigma}$$

158 where,
$$\zeta = 0.3 [\frac{1}{(1-\phi)^{1/3}} - 1] \operatorname{Re}^{1/2} \operatorname{Pr}^{1/3}$$

Presented in Figure 1 are also the curves of Nusselt number versus Reynolds numbers for different porosities based on the model of Nelson and Galloway [22]. It is worth noting that the system of one sphere grain in a fluid is assumed to have the porosity of 1. The Nelson and Galloway curves of Figure 1 therefore represent natural sediments at lower to intermediate porosities and at a porosity of 1 the extreme case of heat transfer between fluid and a single sphere.

165

166 **2.2 Forward two-domain numerical model**

Both the proposed correlation based on the Kunii and Smith [28] data and Nelson and Galloway
[22] theory were embedded into a finite element numerical code to forward model the twotemperature equations (1 & 2) for physical parameters typical of streambed materials [11] (also
shown in Table 1). In the analysis, *Pe* was varied by changing the pore water velocity (~0.01,

171 0.04, 0.09 and 0.3 m/d) and solid thermal conductivity (the upper and lower bound of thermal 172 conductivity of solids are $k_{s_min} = 0.8 \text{ W}(\text{mC})^{-1}$ and $k_{s_max} = 2.5 \text{ W}(\text{mC})^{-1}$) [35]. While we 173 recognise that this velocity range is at the lower end for typical streambeds, using realistic 174 thermal properties it is as high a range as is possible while staying within the *Re* range of the 175 Kunii and Smith [28] data on which our heat transfer correlation is based.

For a particular combination of parameters, equations 8 & 9 were solved for Nu and then h_{sf} was 176 extracted from equation 5 and used in equations 1 & 2. In order to solve Eqs. 1 and 2 177 simultaneously, the initial fluid temperature was used to calculate the solid temperature with the 178 obtained heat transfer coefficient. The obtained solid temperature is then used to calculate new 179 fluid temperature. The ith-step fluid temperature was then compared with i-1th step fluid 180 temperature using a least square technique to check the convergence. The convergence is 181 considered satisfied for a temperature error of 0.01 °C. A two dimensional mesh with 21 nodes 182 183 along x-axis (0.1 m) and 8421 nodes along y axis (4.0 m) with 10 mins time steps were used in the numerical simulation. The depth of 4 m to the lower boundary was sufficient to not influence 184 the results extracted from the upper 0.45 m used for the analysis. 185

Standard Galerkin and Characteristic Galerkin Finite Element discretization techniques [36, 37] with a least square method were used to simultaneously solve for solid and fluid temperatures (equations 1 & 2). Natural heat convection due to buoyancy effects was neglected assuming that the forced convection dominates the heat transfer process [17]. It is also noteworthy that, for the range of *Re* investigated in this study, the thermal dispersion was negligible [19].

Since most studies of groundwater-surface water interactions using heat as a tracer focus on diel 191 temperature signals, we used a daily sinusoidal upper temperature boundary condition for all 192 model scenarios on top and a constant temperature boundary condition (25°C) at the bottom and 193 no flow boundaries at the sides. The initial temperature across the whole model domain was 194 25°C. An amplitude of 4°C for the top boundary starting at 25°C (i.e. $T_0 = T_{ave}$) was used for all 195 runs except for one case where sensitivity to the amplitude was tested. A steady state downward 196 197 fluid flow was assumed and basic physical parameters typical of streambed materials [11] were 198 used. Fluid velocity was varied across a range typically found in the streambed environment for 0.001 < Re < 0.01. However the heat transfer coefficient used for the analysis was not extrapolated 199 200 lower than the lower end of *Re* numbers from the Kunii and Smith [28] experimental data. This

201 prevents from extracting a superficial magnitude for heat transfer coefficient at very low

202 Reynolds numbers (Re<0.001). Models were run for 100 days and the output from the last day

203 of each run was analysed. The finite element numerical discretization of the governing equations

- (1 & 2) is presented in Appendix B.
- 205

206 **2.3 Inverse single-domain analytical model**

207 The output from the two-domain forward models was used as 'synthetic field data' and the 208 amplitude ratios (AR) and phase shifts (PS) of the temperature signal with depth were calculated 209 relative to the upper temperature boundary condition. In a theoretical sense, the fluid and solid temperatures define the upper and lower range of temperature that probes might monitor in 210 streambeds depending on the relative size of the temperature monitoring device and the grain 211 size of the streambed material. In reality, temperature probes will integrate temperature 212 213 responses from the fluid and solid. However for this analysis, rather than choosing an arbitrary averaging of temperatures which would be site-dependent varying with the type of field 214 instrument used and streambed material, we inverted the data for the fluid and solid separately 215 to show the maximum differences that could arise. Therefore, to represent this range within the 216 synthetic data derived from the forward models, ARs and PSs were calculated for the individual 217 temperatures of the fluid (T_f) and solid (T_s) phases throughout the analysis. The AR and PS 218 219 values were then inverted using the commonly used equation which assumes LTE [6] via the 220 equations proposed by Hatch et al. [8] (and the 'known' porosity and thermal parameters in the 221 forward model) to produce values of pore water velocity at depths of 0.1, 0.2, 0.3, 0.35, 0.4 and 0.45 m. Errors in fluid velocity were calculated by comparing the inverse model results with 222 those used in the forward models. For the inversions the bulk thermal diffusivity, D, was 223 224 assumed to be given by the following average of the solid and fluid phases:

225

$$D_{avg} = \frac{(1-\phi)k_s + \phi k_f}{(1-\phi)(\rho c_p)_s + \phi(\rho c_p)_f}$$

- 226 227
- In this bulk averaging, the fluid and solid phases are considered as parallel resistors allowing the calculation of the overall energy flux through the system.
- 228
- 229 **3. Results and Discussion**

230 **3.1** Heat transfer coefficients for low Reynolds numbers

(10)

The best fit correlation of equation (8) to the Kunii and Smith [28] data takes the form:

232
$$Nu = 2.4 \times 10^{-5} + 285.6 \,\mathrm{Pr}^{-3} \,\mathrm{Re}^{2.7}$$

The correlation is shown against the data in Figure 1 alongside output from the Nelson-233 234 Galloway Model (NGM). For the modelled porosity of 0.3 used here, the agreement between the Kunii and Smith Correlation (KSC) and the NGM is good for practical applications at 235 Re=0.01 where the ranges of applicability overlap. This gives confidence in the approach taken 236 here for estimating the heat transfer coefficient. Note that the curves shown for the highest 237 porosities are unrealistic for natural materials but can be realistic for heat transfer within loosely 238 239 packed beds used in chemical reactors. One sphere grain is assumed to have a porosity of 1 and therefore the curves with higher porosity approach the case of heat transfer between fluid and a 240 single sphere. 241

242

3.2 Simulated local thermal non-equilibrium between solid and fluid phases for sinusoidal varying temperature input

Marked differences, up to approximately 1 °C in the modelled cases, were found between the 245 solid and fluid phase temperatures derived from the two-domain model at a range of depths and 246 Pe (and Re) with a surface temperature amplitude of 4 °C and solid thermal conductivity of 247 either 0.8 or 2.5 $W(mC)^{-1}$. Figure 2 illustrates this for a depth of 0.2 m and for high and low Re 248 $(2.5 \times 10^{-4} \text{ and } 7.5 \times 10^{-3})$. The figure also includes the case with thermal equilibrium (e.g. the 249 Hatch equation [8]) and the purely conductive case for comparison. At the low Re of 2.5×10^{-4} 250 the purely conductive case and the LTE case are producing almost identical temperature 251 fluctuations at 0.2 m depth. This illustrates that for this low Re identifying a velocity different 252 253 from zero probably leads to inaccuracy. However, for the two-domain model the temperature 254 fluctuations for solid and fluid differ from each other as well as from the conductive and the 255 LTE cases (both in terms of amplitude and phase). It is interesting to note that the temperature fluctuations for the solid and fluid cannot be combined (by some weighed average) to produce 256 the one-domain analytical LTE temperature fluctuations since they are both simultaneously 257 lower (or both higher) than the LTE temperature. At higher Re (=7.5×10⁻³), there is now a 258 distinct difference between the conductive case and the LTE case. However, the temperature of 259 260 fluid and solid from the two-domain model and LTE case are almost identical for high and low

solid thermal conductivities showing that the two-domain system is approaching thermal equilibrium.

We extracted the difference between the sinusoid amplitude of the solid and fluid temperatures 263 264 (ATD) as a measure of the thermal disequilibrium. In order to investigate the effect of change in amplitude of surface temperature on ATD at different Reynolds numbers, four temperature 265 sinusoids with amplitude of 1, 2, 3 and 4°C were applied on the surface boundary (Figure 3) 266 and the response was measured at 0.2 m depth. In this analysis, the volumetric heat transfer 267 coefficient ($h_{sf.}a_{sf}$ in equation 2) was set constant (200 W(m³C)⁻¹, Re = 0.0056) in order to 268 analyse only the effect of velocity on ATD (and neglect the effect of heat transfer coefficient). 269 Figure 3 indicates that the lower the temperature amplitude applied at the top boundary the 270 lower the resultant ATD. Moreover, the increase in velocity gives rise to increasing values for 271 ATD particularly when it passes the threshold of Pe = 0.0074 (or Re=0.01). This is due to the 272 273 fact that an increase in velocity leads to a higher localised temperature gradient at the grain boundary; greater thermal non-equilibrium occurs in these modelled conditions as conduction 274 into the grains cannot keep pace with the advective flux of heat through the fluid (i.e. higher 275 grain Pe). 276

277

278 **3.3** Error in derived streambed fluid velocity when assuming local thermal equilibrium

279 The relative $\left(\frac{v_{ARorPS} - v_{actual}}{v_{actual}}\right)$ and absolute $\left(v_{ARorPS} - v_{actual}\right)$ errors in pore water velocity (from

both the AR and PS [8]) using T_s , or T_f as a function of *Pe* are presented in Figure 4a-d. From 280 281 Fig. 4, the errors in derived velocity estimates converge to zero value for all cases as Pe 282 increases whether using T_s , or T_f except the PS velocity errors obtained from T_f and high solid thermal conductivity (k_{high}) . So, while the increase in advective flux (Pe) tends to thermally 283 284 disequilibrate the system (Fig. 3), this is more than compensated by an increased heat transfer 285 coefficient (h_{sf}) at higher velocities which tends to increase equilibrium between phases, leading 286 to more equilibrium at higher Pe(Re) in the range considered here (This is summarised 287 conceptually in Fig. 7).

It can be seen from Fig. 4 that the *AR* derived relative and absolute velocity errors are negative and decrease with depth using T_f and high solid thermal conductivity (k_{high}) at low *Pe* (low *Re*), whereas the errors are positive using T_s with the same k_{high} and at low *Pe*. This is attributed to

the fact that *AR* values of the solid and fluid phases are different to that of the local thermal equilibrium case (i.e. *AR* derived from the 1-D analytical solution based on the LTE assumption). In order to compare the *AR* values of the numerical analysis to that of the analytical solution at different *Re* (= 2.5×10^{-4} and 7.5×10^{-4}) **Fig. 5** is presented (it should be noted that *Pe* is replaced with *Re* in Fig. 5 due to the fact that *Pe* varies with change in solid thermal conductivity).

297

As an example, the AR values of the solid phase, with high solid thermal conductivity (k_{high}) at 298 low Pe (Re= 2.5×10^{-4}), are higher than that of the local thermal equilibrium case leading to 299 higher derived velocities than for the LTE case and thus positive errors. AR values of the fluid 300 phase, with high solid thermal conductivity (k_{high}) at low Pe (Re=2.5×10⁻⁴), are lower leading to 301 lower velocities than the LTE case and thus negative errors. It can also be seen from Fig. 2 that 302 303 the temperature fluctuations of the LTE case is lower than the temperature fluctuations of the solid phase with k_{high} and higher than the temperature fluctuations of the fluid at low Pe 304 $(Re=2.5\times10^{-4})$. The physical basis for these deviations is that at low Pe, the heat exchange 305 between phases becomes inefficient and therefore, using k_{high} , the heat transport in the solid 306 phase becomes much quicker than that within the fluid. 307

308

Using a lower solid thermal conductivity (k_{low}) and low Pe (Re=2.5×10⁻⁴), on the other hand, the 309 ARs using either T_s or T_f are both greater than those for the local thermal equilibrium case and 310 therefore positive velocity errors are obtained. Again it can be explained by the fact that at low 311 *Pe* the heat exchange between phases is inefficient and since the solid thermal conductivity is 312 low (very close to fluid thermal conductivity) the solid and fluid phases end up behaving 313 314 similarly. The reason why the AR value of the LTE case is slightly lower than both the solid and fluid ARs is because of the difference in the thermal diffusivity of each phase and that of the 315 LTE case. Although the thermal conductivity of the LTE case sits between the solid and fluid 316 thermal conductivities, its thermal diffusivity may sit between or below the solid and thermal 317 318 phases due to a different volumetric heat capacity. And because the thermal diffusivity affects 319 the rate of heat transfer, lower magnitude of AR is observed compared to that of solid and fluid (where the thermal diffusivity of LTE case sits below the solid and thermal phases). It is 320 noteworthy that the diffusivity is the function of both the thermal conductivity and the 321

- volumetric heat capacity. When moving toward higher Pe ($Re=7.5\times10^{-4}$), the error approaches zero showing that the system reaches local thermal equilibrium.
- 324

The relative errors in the PS derived velocity estimates (Fig. 4) have similar trends and greater 325 magnitudes compared to those derived using ARs especially at lower end of Pe. From Fig. 4d, it 326 can be seen that the PS derived absolute velocity errors stay constant at relatively lower 327 velocities (Pe). Thus the relative errors increase only due to a reduction in the actual pore water 328 329 velocity. Due to the fact that the AR and PS methods are sensitive to different velocities [8], the PS method loses its sensitivity at lower range of velocity and the same velocity estimate is 330 returned. In addition, the errors at higher velocities do not converge to the absolute zero which is 331 resulted from the effect of local thermal non-equilibrium on the phase shift of the temperature 332 data. The obtained PS values of the numerical analysis and analytical solution at different Re 333 $(=2.5\times10^{-4} \text{ and } 7.5\times10^{-4})$ are also presented in **Fig. 6** for comparison. 334

335

Since the errors we have reported here are significant, especially for relatively low Pe (relative 336 errors up to 30 and 150 are obtained from AR and PS), we have compared the parameter range 337 of our results to laboratory studies which present data with which it is possible to assess the 338 robustness of a single-domain equation (implicitly assuming the validity of LTE) in deriving 339 340 stream bed velocities using diurnal temperature signals. Surprisingly, given the ever increasing number of field applications using such an approach there are, to our knowledge, only 3 341 laboratory studies of relevance. Rau et al. [19] found generally good agreement between 342 experimental and theoretical expectations in a study conducted at a range of Re above the data 343 presented here, in the range where we would expect the LTE assumption to be valid. Munz et 344 345 al. [38] and Lautz [39] present results which may cross over with the range of Re we have analysed here although, unfortunately, neither paper is explicit regarding the grain size 346 distribution used in their experiments. However, using a typical range of grain sizes for fine 347 sand [39] and medium sand [38] the minimum Re studied may have been approximately $6 \cdot 10^{-3}$ 348 and $2.5 \cdot 10^{-3}$ respectively which are within the range of values where we would expect LTE to 349 350 breakdown. In the Lautz [39] experiments, we note that significant discrepancies were found between velocities derived using AR and PS, which remain unexplained and that might be due to 351 LTNE, although other effects such as heterogeneity can also induce such discrepancies [40, 41]. 352

- In the Munz et al. [38] experiments, increasing discrepancies are apparent between the measured and modelled flow velocities as the flow rate decreases. These observations are consistent with the understanding of LTNE described in this paper, and we propose that false assumptions of LTE may have contributed to these reported errors.
- The errors that could arise due to a false assumption of LTE may be of the same order of magnitude as errors due to other factors such as non-uniform flow fields [10, 11], sediment heterogeneity [12], measurement error and difficulties in estimating thermal parameters [13, 14].
- 360

361 **4. Conclusion**

Despite a large body of literature describing the fundamentals of heat transfer in porous media, the plethora of studies which have applied heat as a tracer in streambeds have, to our knowledge without exception, assumed local thermal equilibrium between solid and fluid phases. However, there is evidence from existing theory and empirical evidence that this assumption may not be valid in many instances [22, 28].

Here we have derived a correlation for the heat transfer coefficient at low *Re* using well known 367 experimental data (KSC) which is in good agreement with a physically based model (NGM). 368 Our analysis reveals that two main mechanisms control the degree of thermal equilibrium 369 between the solid and fluid phases in a typical streambed: the ratio of the conductive to 370 371 advective heat transport (described by the grain thermal Pe) and the heat transfer coefficient which is related to the Re (Figure 7). These processes act against each other; higher advection 372 tends towards disequilibrium between phases while at high velocities this process is more than 373 outweighed by an increasing heat transfer coefficient which tends to move the system towards 374 equilibrium. Including these processes in a two-domain heat transport model we have shown 375 376 that the LTE assumption may break down at *Re*<0.01 for typical streambed thermal parameters. Furthermore, this model output was then inverted using a 1D analytical model which assumes 377 378 LTE, to show that considerable relative errors in streambed velocity estimates may result at low Re (or Pe) if the possibility of LTNE is ignored. In general, these errors are higher at relatively 379 380 lower Re and may lead to significant inferred flows from data inversions based on the LTE 381 assumption (0.3 m/d using AR and 1.3 m/d using PS) when in fact the real flow is small or zero. Such errors may be of the same order of magnitude as other known uncertainties in streambed 382 heat tracing [10-14]. 383

These results have important implications for interpreting and predicting streambed temperature dynamics, critical for improving the understanding of controls on stream ecology and biogeochemical processes. More laboratory studies are urgently needed to supplement the sparse existing data in this area and further test the findings of this study. In particular, the data and models on which this study is based was for homogeneous media and diel temperature signals, and it is to be expected that results will significantly differ for real field conditions; such data are required to enable a more complete physical understanding of heat transport processes in real streambeds to be derived.

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415 Appendix A

- **Table A.** Details of fitting parameters to the experimental data of Kunii and Smith [1961] in
- 418 Figure 1 using DATAFIT software.

Results from project "LTNE"		
	$Nu/Pr^2 = a+b*Re^c$	
Model Definition:	Where $a = \alpha / Pr^2$, $b =$	
	$K_1 \times Pr^{(1/3)}/Pr^2$ and $c=q$	
Number of observations	41	
Number of missing observations	0	
Solver type	Nonlinear	
Nonlinear iteration limit	250	
Diverging nonlinear iteration limit	10	
Number of nonlinear iterations performed	61	
Residual tolerance	1.00E-10	
Sum of Residuals	9.31E-15	
Average Residual	2.27E-16	
Residual Sum of Squares (Absolute)	3.63E-11	
Residual Sum of Squares (Relative)	3.63E-11	
Standard Error of the Estimate	9.78E-07	
Coefficient of Multiple Determination (R^2)	8.37E-01	
Proportion of Variance Explained	83.68%	
Adjusted coefficient of multiple determination (Ra^2)	0.83	
Durbin-Watson statistic	1.53	

425 Table B. Regression variable results for the experimental data of Kunii and Smith [1961]

426 including the best fit and confidence intervals of 68%, 90%, 95% and 99% from DATAFIT

427 software.

Variable	Value	Standard	t-ratio	Prob(t)
Α	7.35E-07	4.48E-07	1.640975375	0.10906
В	15.3962065	42.61194092	0.361312021	0.71987
С	2.687445266	0.51686944	5.199466357	0.00001
68% Confidence				
Variable	Value	68% (+/-)	Lower	Upper
Α	7.35E-07	4.51E-07	2.84E-07	1.19E-06
В	15.3962065	42.93579167		58.33199817
С	2.687445266	0.520797648	2.166647618	3.208242914
90% Confidence				
Variable	Value	90% (+/-)	Lower	Upper
Α	7.35E-07	7.55E-07	-2.02E-08	1.49E-06
В	15.3962065	71.84373239	-	87.23993889
С	2.687445266	0.871441876	1.816003389	3.558887142
95% Confidence				
Variable	Value	95% (+/-)	Lower	Upper
Α	7.35E-07	9.07E-07	-1.72E-07	1.64E-06
В	15.3962065	86.2636132	-70.8674067	101.6598197
С	2.687445266	1.046350495	1.641094771	3.73379576
99% Confidence				
Variable	Value	99% (+/-)	Lower	Upper
Α	7.35E-07	1.21E-06	-4.80E-07	1.95E-06
В	15.3962065	115.5422778	-	130.9384843
C	2.687445266	1.401491487	1.285953778	4.088936753
Variance Analysis				
Source	DF	Sum of	Mean	F Ratio
Regression	2	1.86E-10	9.32E-11	97.42871476
Error	38	3.63E-11	9.56E-13	
— 1				

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433 Appendix B

Numerical discretization: the standard and Characteristic Galerkin techniques are used to
discretize the governing equations of the two-domain heat transport problem (equations 1 and
2). It results in the following system of equations for a two dimensional problem:

437
$$[-(M + \Delta t[H - M_3])][\Delta T_i^s] = [\Delta t[H - M_3]T_i^s(t_{i-1}) - \Delta tM_3T_i^f(t_i)]$$

438
$$[-(M + \Delta t(C - K_1 - K_2) - \Delta tM_2)][\Delta T_i^f] = [\Delta t[(C - K_1 - K_2) - M_2]T_i^f(t_{i-1}) + \Delta tM_2T_i^s(t_i)]$$
438

440 where *i* is the time step; \vec{T} is the temperature vector; $\vec{T}_T = (T_1 \ T_2 \ \dots \ T_n)$; *T* is the nodal 441 temperature; subscripts s and f represent the solid and fluid phases respectively; Δt represents 442 the time increment and the matrices are defined as:

443
$$\vec{\tilde{M}} = \int_{V_e} [N_T]^T [N_T] dV$$

$$444 \qquad \vec{\vec{C}} = \int_{V_e} [N_T]^T \frac{\partial}{\partial x} \left(\frac{\langle k \rangle^f}{(\rho c_p)_f} + f\left(\vec{\vec{\beta}}, \vec{v}_x\right) \right) \frac{\partial [N_T]}{\partial x} \left\{ \chi \right\}^n dV + \int_{V_e} [N_T]^T \frac{\partial}{\partial y} \left(\frac{\langle k \rangle^f}{(\rho c_p)_f} + f\left(\vec{\vec{\beta}}, \vec{v}_y\right) \right) \frac{\partial [N_T]}{\partial y} \left\{ \chi \right\}^n dV + V_{V_e} [N_T]^T \frac{\partial}{\partial y} \left(\frac{\langle k \rangle^f}{(\rho c_p)_f} + f\left(\vec{\vec{\beta}}, \vec{v}_y\right) \right) \frac{\partial [N_T]}{\partial y} \left\{ \chi \right\}^n dV + V_{V_e} [N_T]^T \frac{\partial [N_T]}{\partial y} \left\{ \chi \right\}^n dV$$

446
$$\vec{K}_{2} = \frac{\Delta t}{2} v_{x}^{d} \int_{V_{e}} \left[\frac{\partial}{\partial x} (v_{x}^{d} \frac{\partial [N_{T}]}{\partial x} \{\chi\}^{n} + v_{y}^{d} \frac{\partial [N_{T}]}{\partial y} \{\chi\}^{n}) \right] dV + \frac{\Delta t}{2} v_{y}^{d} \int_{V_{e}} \left[\frac{\partial}{\partial y} (v_{x}^{d} \frac{\partial [N_{T}]}{\partial x} \{\chi\}^{n} + v_{y}^{d} \frac{\partial [N_{T}]}{\partial y} \{\chi\}^{n}) \right] dV$$

447
$$\vec{M}_2 = \int_{V_e} \frac{h_{sf} a_{sf}}{\phi(\rho c_p)_f} [N_T]^T [N_T] dV$$

448
$$\vec{M}_{3} = \int_{V_{e}} \frac{h_{sf} a_{sf}}{(1-\phi)(\rho c_{p})_{s}} [N_{T}]^{T} [N_{T}] dV$$

449
$$\vec{H} = \int_{V_e} [N_T]^T \frac{\partial}{\partial x} (\frac{\langle k \rangle^s}{(1-\phi)(\rho c_p)_s}) \frac{\partial [N_T]}{\partial x} (\chi)^n dV + \int_{V_e} [N_T]^T \frac{\partial}{\partial y} (\frac{\langle k \rangle^s}{(1-\phi)(\rho c_p)_s}) \frac{\partial [N_T]}{\partial y} (\chi)^n dV$$

450 where N_T is the finite element shape function of temperature, V is the spatial area of an element 451 and χ is the variable.

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Figure 1 Variation of *Nu* with *Re*: Kunii & Smith (1961) experimental data alongside our correlation and the Nelson Galloway Model (NGM) results for a variety of porous material porosities (ϕ).

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Figure 2 Sinusoidal temperature fluctuations at the surface and at 0.2 m depth for Re numbers of 7.5×10^{-3} and 2.5×10^{-4} and high (2.4 W(mC)⁻¹) and low (0.8 W(mC)⁻¹) solid thermal conductivities. Temperatures were calculated for the solid and the fluid by the two-domain model (as outlined in the methodology), and for the assumption of local thermal equilibrium (LTE) using the method by Hatch et al. [8] and for the case of no flow (thermal diffusion only).

Figure 3. the amplitude of the temperature difference (ATD) as a function of Re at four different temperature amplitudes (1, 2, 3 and 4 °C) at the stream-sediment temperature boundary condition. For this simulation the heat transfer coefficient has been held constant and the depth of measurement is 0.2 m.

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Figure 4. a) *AR* derived relative velocity error, b) *PS* derived relative velocity error c) *AR* derived absolute velocity error and d) *PS* derived absolute velocity error vs Pe_{avg} using solid and fluid phase temperatures and higher and lower values of solid thermal conductivity ($k_{s_min}=0.8$ W(mC)⁻¹ and $k_{s_max}=2.4$ W(mC)⁻¹). The velocity range is ~0.01-0.3 m/d. For all plots the set of curves for each symbol represents velocity error estimates for depths of 0.1, 0.2, 0.3, 0.35, 0.4 and 0.45 m.

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Figure 5. The amplitude ratio (*AR*) of the temperature signal vs depth at high (= 7.5×10^{-3}) and low (= 2.5×10^{-4}) Reynolds numbers for high and low solid thermal conductivities using solid and fluid temperatures. Also shown are the *ARs* derived using the 1-D analytical solution which assumes LTE [Hatch et al, 2006].

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Figure 6. The phase shift (*PS*) of the temperature signal (*PS*) vs depth at high (= 7.5×10^{-3}) and low (= 2.5×10^{-4}) Reynolds numbers for high and low solid thermal conductivities using solid and fluid temperatures. Also shown are the *PS* derived using the 1-D analytical solution which assumes LTE [Hatch et al, 2006].

Figure 7. The relative importance of advective heat transport through the fluid, and heat transfer between the solid and the fluid phases at high and low Re. a) At low flow rates the heat transfer is relatively inefficient at thermally equilibrating the solid and fluid phases and LNTE is possible. b) At high rates of fluid advection (high *Pe*) even though heat is advected fast through the porous media the heat transfer is far more efficient and helps maintain LTE.
Table 1. Physical data used in the study.











Depth (m)



Depth (m)



Table 1 637

	Parameter	Unit	Symbol	Value
	Solid Thermal Conductivity	$W(mC)^{-1}$	k_{s_\min} & k_{s_\min}	0.8 & 2.5
	Water Thermal Conductivity	W(mC) ⁻¹	$k_{_f}$	0.58
	Water Specific Heat Capacity	J(kgC) ⁻¹	<i>c</i> _{<i>f</i>}	4183
	Solid Specific Heat Capacity	J(kgC) ⁻¹	c _s	750
	Water Density	kg m ⁻³	<i>P</i> _j	999.7
	Solid Density	kg m ⁻³	ρ_s	2650
	Porosity	-	φ	0.3
	Longitudinal Thermal Dispersivity	s	β_{i}	1.478
	Transverse Thermal Dispersivity	S	β_{i}	0.4

640 Highlights

641	• We have derived a correlation for heat transfer coefficient at low <i>Re</i>
642	• Local thermal equilibrium may not be a valid assumption in sediments' heat transfer
643	Error in temperature derived velocity estimates may be obtained using LTE
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