Humans Adapt Multi-Objective Control of Stepping To Perform Lateral Maneuvers While Walking

Short Title: Adapting Stepping to Laterally Maneuver

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ABSTRACT [294 Words – Max = 300]
To successfully traverse their environment, humans must often perform maneuvers to achieve desired task goals while simultaneously maintaining balance. Humans accomplish these tasks primarily by modulating their foot placements. As humans are more unstable laterally, we must better understand how humans modulate lateral foot placement. We previously developed a theoretical framework and corresponding computational models to describe how humans regulate lateral stepping during straight-ahead continuous walking. This framework yields goal functions for step width and lateral body position that define the walking task and determine the set of all possible task solutions as Goal Equivalent Manifolds (GEMs). Here, we used this framework to determine if humans can regulate lateral stepping during non-steady-state lateral maneuvers by minimizing errors in accordance with these goal functions. Twenty young healthy adults each performed four lateral lane-change maneuvers in a virtual reality environment. Within our general lateral stepping regulation framework, we first re-examined the requirements of such transient walking tasks. Doing so yielded new theoretical predictions regarding how steps during any such maneuver should be regulated to minimize error costs, consistent with the goals required at each step and with how these costs are adapted at each step during the maneuver. Humans performed the experimental lateral maneuvers in a manner consistent with our theoretical predictions. Furthermore, their stepping behavior was well modeled by merely adapting the parameters of our previous lateral stepping models from step to step. To our knowledge, our results are the first to demonstrate humans can use evolving cost landscapes in real time to perform such an adaptive motor task and, furthermore, that such adaptation can occur quickly – over only one step. Thus, the predictive capabilities of our general stepping regulation framework extend to a much greater range of walking tasks beyond just normal, straight-ahead walking.
AUTHOR SUMMARY [167 Words – Range = 150-200]

When we walk in the real world, we rarely walk continuously in a straight line. Indeed, we regularly have to perform other tasks like stepping aside to avoid an obstacle in our path (either fixed or moving, like another person coming towards us). While we have to be highly maneuverable to accomplish such tasks, we must also remain stable to avoid falling while doing so. This is challenging because walking humans are inherently more unstable side-to-side. Sideways falls are particularly dangerous for older adults as they can lead to hip fractures. Here, we establish a theoretical basis for how people can accomplish such maneuvers. We show that humans execute a simple lateral lane-change maneuver consistent with our theoretical predictions. Importantly, we show that they can do so using simple adaptations of the same step-to-step regulation strategies they use to walk in a straight line. Moreover, these same control processes also explain how humans trade-off side-to-side stability to gain the maneuverability they need to perform such lateral maneuvers.
INTRODUCTION

To successfully traverse our environment, we humans must adapt to a wide variety of environment contexts and changing task goals (e.g., Fig. 1A), all while maintaining balance. Indeed, humans readily avoid obstacles [2] and/or other humans [3], step to targets [4-6], move laterally [7-9], navigate complex terrain [10-12], and respond to destabilizing perturbations [13, 14]. To do so requires a high degree of maneuverability [7] and, potentially, the ability to trade-off stability for maneuverability [8, 9]. While maneuverability has been widely studied in animal locomotion [15, 16] and robotics [16, 17], far less is known about how humans accomplish such tasks.

Humans accomplish these sorts of adaptive walking tasks primarily by modulating their foot placements. Appropriate foot placement at each step can redirect center-of-mass accelerations and thus allows humans to maintain balance [18-21]. This is especially important in the lateral direction where humans are thought to be more unstable [19, 22, 23]. Indeed, laterally-directed falls are particularly injurious in older adults [24, 25]. Thus, it is important to better understand how humans adapt their lateral foot placements to maintain balance while simultaneously achieving walking task goals.

Human walking exhibits considerable variability [26, 27], redundancy (i.e., the body has more mechanical degrees of freedom than necessary to perform any movement) [28], and equifinality (i.e., humans can perform most tasks an infinite number of ways) [29-31]. Computational models are necessary to fully understand how humans perform accurate, goal-directed walking movements in the context of these challenges. Most prior models of human walking (e.g., [22, 23, 32, 33]) address within-step “control,” defined here as the processes that drive the dynamics of each step to remain viable (i.e., to prevent falling) [34]. While ensuring viability is indeed necessary to walk [35], it is not sufficient: humans engage in purposeful walking with a destination to reach and/or other tasks to achieve. Control models that “just walk” (i.e., remain viable) do not address how humans achieve such goal-directed walking.

We posit that locomotion is functionally hierarchical, consisting of both within-step control and step-to-step regulation. To compliment within-step control models, we developed a motor regulation framework to determine how humans adjust successive stepping movements [1, 29, 34, 36]. For a given walking task (e.g., Fig. 1A), this framework proposes goal functions that theoretically define the task and, accounting for equifinality, determine the set of all possible task solutions as Goal Equivalent Manifolds (GEMs) [37-40]. The goal functions are incorporated into task-level costs, which we then use in an optimal control formalism [41, 42] to generate relatively simple, phenomenological models of step-to-step motor regulation (i.e., “motor regulation templates” [1, 29, 30]). For any proposed goal function, these templates predict how humans would attempt to drive that goal function to zero at the next step, thus achieving perfect task performance on average. For an appropriate choice of goal functions, these models successfully replicate human step-to-step
dynamics in both the fore-aft [29] and lateral [1] directions. In the lateral direction in particular, multi-objective regulation of primarily step width ($w$) and secondarily lateral position ($z_B$) captures human step-to-step dynamics during continuous straight-ahead walking (Fig. 1B-C) [1, 13, 43]. Whether explicit models are constructed or not, this overall theory and its hierarchical control/regulation schema provides a powerful framework from which to interpret experimental results [13, 31, 43-50].

However, humans rarely perform long bouts of straight-ahead, continuous walking [51, 52]. Instead, humans must frequently perform adaptive locomotor behaviors (i.e., maneuvers) when walking in the real world (e.g., Fig. 1A) [7-9]. Indeed, older adults often fall during such maneuvers because they incorrectly transfer their body weight or they trip [53]. Because both such causes can be prevented with appropriate foot placement [18-21], it is necessary to better understand how humans regulate foot placement as they execute typical lateral maneuvers. Here, we aimed to determine how humans regulate their stepping during a simple lateral maneuver: namely, a single lateral lane-change transition between periods of straight-ahead walking.

It is not clear a priori that our previous lateral stepping regulation framework can also emulate human stepping during lateral maneuvers. This framework was originally developed to model straight-ahead steady-state walking, under assumptions that step-to-step adjustments can be made without changing the fundamental structure of within-step control, and that deviations from perfect performance are small. These assumptions motivated us to select low-dimensional, single-step, linear regulators to model these step-to-step error-
correcting processes [1, 29, 30]. However, walking tasks requiring substantial non-steady-state lateral maneuvers would seem to violate these assumptions. Lateral maneuvers that introduce large deviations from steady-state might, for example, require changes to the within-step control structure, or induce substantial nonlinearity. Lateral maneuvers might also require substantially greater impulses to execute, which may necessitate additional compensatory motor and/or kinematic contributions and could result in higher dimensionality. Furthermore, humans have been shown plan maneuvers more than one step in advance in some contexts [54-57], which suggests the possible need for models that depend on more than one prior step. Thus, one might reasonably posit that humans regulate stepping using entirely different control schemes when executing lateral maneuvers far from steady state.

However, here we demonstrate that humans do indeed regulate lateral stepping during non-steady-state lateral maneuvers in a manner consistent with our previous theoretical framework. We first apply concepts from the general theory in a manner that satisfies the requirements of a non-steady-state lane change maneuver. Specifically, we allow the task goals, task-level costs, and the weighting between competing costs to adapt from each step to the next. We thus derive a model that yields explicit, empirically-testable predictions about how the variability observed during such lateral maneuver tasks will be structured from step to step. We then test these predictions against human experimental data. Lastly, we used the models with these additional adaptive hierarchical elements to simulate how such lateral maneuvers are achieved over a sequence of steps. In doing so, we demonstrate how humans modulate their stepping to trade-off lateral stability for maneuverability to accomplish the lane change task. Thus, our hierarchical motor regulation framework can be applied to a far wider range of walking tasks beyond just straight-ahead, steady-state walking.

RESULTS

Lateral Maneuvers in Humans

We analyzed data from a prior study conducted in our laboratory [58]. Young healthy participants walked in a virtual reality environment. They walked on one of two parallel paths, the centers of which were 0.6m apart, projected onto a 1.2m wide motorized treadmill (Fig. 2A). Following an audible cue, they executed a lateral maneuver from the path they were walking on over to the adjacent path. Importantly, participants were given no explicit instructions about how to execute this maneuver. We analyzed seventy-nine total maneuvers from twenty participants.

Participants variably took 0-to-3 steps between the cue and initiating the maneuver (Fig. 2A[i-ii]). However, once initiated, they performed each maneuver consistently (Fig. 2A[iii]-B). Participants completed nearly all maneuvers with an ipsilateral transition step that involved a single, large step to reach the new path. One participant, however, completed one of their maneuvers by taking a large cross-over step. Additionally, one maneuver from another participant required three steps to reach the new path. These two non-conforming
maneuvers demonstrate that participants had many options: the task itself did not require them to complete the maneuver in any specific way.

Participants completed nearly all lateral maneuvers in 4 non-steady-state steps. Participants first took a
“preparatory” step (step 0) prior to the transition, during which they slightly narrowed their step width and incrementally moved towards the new path. They then took a large “transition” step (step 1) to cover most of the transition distance. Participants took a subsequent “recovery” step (step 2) that again exhibited a slightly narrowed step width. Participants then reached their final new goal position (step 3) and returned to steady-state walking on their new path.

Steady-State Stepping Regulation Cannot Replicate Lateral Transitions

We first used our previous multi-objective model of lateral stepping regulation to test whether humans could regulate stepping during lateral maneuvers using a constant regulation strategy like that used for continuous, straight-ahead walking [1]. This model selects each new foot placement ($z_L$ or $z_R$) as a weighted average of independent predictions that minimize errors with respect to either a constant step width ($w^*$) or lateral position ($z_B^*$) goal, consistent with multi-objective stochastic optimization of error costs with respect to these two quantities (see Methods). For this model, the relative proportion of step width to lateral position regulation was defined by $\rho$, where $\rho = 0$ indicates 100% $z_B$ control (and hence, 100% weight on the $z_B$ cost) and $\rho = 1$ indicates 100% $w$ control (100% weight on the $w$ cost) [1]. This stepping regulation model reproduced the key features of human stepping dynamics during continuous, straight-ahead walking for constant values in the approximate range $0.89 \leq \rho \leq 0.97$ [1].

We assessed whether this model, with any constant value of $\rho$, could emulate the non-steady-state stepping dynamics experimentally observed during the lateral maneuver task. We found that it was not capable of doing so. The model did emulate steady-state stepping dynamics (Fig. 3C; Step -3) for constant values in the range of approximately $0.83 \leq \rho \leq 0.92$, consistent with previous findings [1]. However, no simulations over this range of $\rho$ completed the lateral maneuver in the experimentally observed 4 steps (Fig. 3A-B; red). Furthermore, while model simulations that weighted position and step width regulation similarly (i.e., $\rho \approx 0.5$) approximately emulated average experimentally observed stepping time series and errors (Fig. 3A-B; blue), they failed to emulate the stepping variability humans exhibited during either steady-state (Step -3) or transition (Step 1) steps (Fig. 3C). Indeed, no single constant value of $\rho$ emulated both steady-state and transition stepping dynamics (Fig. 3C).

Re-Thinking Stepping Regulation for Non-Steady-State Tasks

Any biped (human, animal, robot, etc.) must enact step width and/or lateral position regulation via left and right foot placement (Fig. 1C). The stepping goals, [$z_B^*$, $w^*$], which guide steady-state walking, individually form diagonal, orthogonal Goal Equivalent Manifolds (GEMs) when plotted in the [$z_L$, $z_R$] plane (Fig. 4A) [1]. The intersection of these GEMs represents the multi-objective goal to maintain both $z_B^*$ and $w^*$ and therefore defines the foot placement goal, [$z_L^*$, $z_R^*$], for the task. When viewed in this manner, it is evident that nearly any substantive change in either stepping goal (i.e., $\Delta z_B^*$ and/or $\Delta w^*$) will induce a diagonal shift of the
corresponding GEM(s) in the $[z_L, z_R]$ plane (Fig. 4A). This will necessitate corresponding changes in both left and right foot placement, which therefore cannot be accomplished in a single step (Fig. 4A).

At minimum, two consecutive steps are required to execute nearly any maneuver involving some $\Delta z_B^*$ and/or
$\Delta w^*$ (Fig. 4B). The first step must be taken by either the left or right foot to either of two possible intermediate foot placement goals in the $[z_L, z_R]$ plane (Fig. 4B). Indeed, we can derive exact stepping goals for this intermediate step algebraically (see Methods). However, equifinality exists in both the number and placements of steps that can be used to accomplish any $\Delta z_B^*$ and/or $\Delta w^*$ maneuver. For example, we experimentally observed a four-step strategy to be most typical (Fig. 2). To represent such a strategy, we incorporated (see Methods) a lateral offset ($\varepsilon$) for the preparatory and recovery steps and determined the foot placements, $[z_L, z_R]$, and corresponding to transition stepping goals, $[z_B^*, w^*]$, at each of the corresponding
preparatory, transition, and recovery steps (Fig. 4C).

However, specifying these foot placements alone does not capture how any given biped might perform these steps. Along any GEM, deviations tangent to the GEM are “goal equivalent” because they do not introduce errors with respect to the goal. Conversely, deviations perpendicular to the GEM are “goal relevant” because they do introduce such errors [29]. Humans typically exhibit greater variability along GEMs they exploit [37, 41, 42]. For multi-objective lateral stepping regulation, the $z^*_B$ and $w^*$ GEMs are orthogonal (Fig. 4A). Thus, goal equivalent deviations with respect to either GEM are goal relevant with respect to the other (Fig. 4D) [1]. Hence, the ratio of the $\delta_{z_B}$ and $\delta_w$ deviations with respect to both the $z^*_B$ and $w^*$ GEMs theoretically reflects the relative weighting of $z_B$ and $w$ regulation. For example, during steady-state walking, the distribution of human steps is anisotropic: that is, the steps are strongly aligned along the $w^*$ GEM (such that $\delta_{z_B}/\delta_w >> 1$) because humans heavily weight step width over position regulation [1].

When the stepping goals change (i.e., $\Delta z^*_B$ and/or $\Delta w^*$), our theoretical framework thus posits empirically testable hypotheses about how these $\delta_{z_B}/\delta_w$ ratios should also change at each step. For a minimum two-step strategy (Fig. 4B), foot placement at the intermediate step should be as accurate as possible to minimize stepping errors at the final step. Theoretically then, our framework predicts that stepping distributions at this ideal intermediate step should not exhibit equifinality and hence would be approximately isotropic (i.e., circular), reflecting equal weighting of step width and position regulation (Fig. 4E). However, taking more intermediate steps during any such $\Delta z^*_B$ and/or $\Delta w^*$ maneuver reduces the accuracy required at each individual step, as errors can be corrected at subsequent intermediate step(s) prior to reaching the final goal. Our theory thus predicts that stepping distributions for multiple intermediate steps should be more isotropic than at steady-state, but perhaps less than the ideal two-step case. For example, for a four-step strategy with a large primary transition step and smaller preparatory and recovery steps (Fig. 4C), stepping distributions are predicted to be most isotropic at the primary transition step and intermediately isotropic at the preparatory and recovery steps (Fig. 4F).

We thus hypothesized that humans would regulate their stepping movements across these lane-change maneuvers in accordance with these theoretical predictions. We tested this hypothesis by plotting experimental data in the [$z_L$, $z_R$] plane and fitting 95% prediction ellipses to the data for each relevant step (Fig. 5). During steady-state walking both before and after the maneuver, participants’ steps were strongly aligned to the constant-$w^*$ GEM (Fig. 5A), consistent with the expected strong prioritization of step width regulation [1]. During each of the preparatory, transition, and recovery steps, the locations of the experimental stepping distributions were consistent with the intersections of the predicted $z^*_B$ and $w^*$ GEMs at each step (Fig. 5B). Furthermore, the experimental stepping distributions themselves (Fig. 5C) were qualitatively consistent with our theoretical predictions (Fig. 4F). The fitted 95% prediction ellipses at the preparatory and
transition steps were significantly more isotropic than that observed during steady-state walking (Fig. 5C; top), with the primary transition step being most isotropic as theoretically predicted (Fig. 4F). Thus, these experimental data strongly support our hypothesis that humans regulate their movements from one step to the next during these lane-change maneuvers by adapting their stepping goals and the relative weighting of $z_B$ and $w$ regulation.

Adaptive Stepping Regulation for Lateral Maneuvers

Our theoretical predictions (Fig. 4) and confirmatory experimental results (Fig. 5) both strongly suggest that humans can accomplish a range of substantially non-steady-state lateral maneuvers by simply changing how they weigh different task-level costs from each step to the next. They do not need to adopt some entirely different stepping regulation process. We therefore sought to determine the extent to which our previous multi-objective model [1] could emulate such variations in stepping dynamics if we allowed the parameters of our model to adapt from step-to-step.

Our regulator models are linear update equations (see Methods). Their behavior is controlled primarily by three basic parameters: a pair of target values or stepping goals, $[z_B^*, w^*]$, a weight that indicates each regulator’s relative importance, $\rho$, and the additive noise, $\sigma_a$, that represents the strength of physiological perceptual/motor noise. Here, we conducted three sequential numerical experiments to assess the effects of...
adapting each of these model parameters on key stepping dynamics: time series, errors, and variance distributions. We first incorporated adaptive stepping goals that we derived theoretically to approximate an idealized four-step transition strategy (Fig. 4C). Next, we adapted the proportionality parameter, $\rho$, to reflect the predicted stepping distributions (Fig. 4F). Finally, we increased the additive noise to reflect the observed increase in variability during the lateral maneuver. Importantly, we did not attempt to precisely estimate the model parameters, but rather selected parameters based on the idealized theoretical considerations described above. This approach is analogous to using “templates” of legged locomotion to reveal the basic principles of human walking by testing fundamental hypotheses about the underlying regulation strategies [59, 60].

We algebraically derived (see Methods) new stepping goals, $[z_B^*, w^*]$, for each consecutive step (Fig. 6A) that approximated an idealized four-step lane-change maneuver strategy (Fig. 4C). These derived stepping goals included a slightly narrower $w^*$ and slight shift in $z_B^*$ towards the new path at the preparatory step, a much wider $w^*$ and large shift in $z_B^*$ towards the new path at the transition step, and a slightly narrower $w^*$ and slight shift in $z_B^*$ towards the new path again at the recovery step. All other model parameters were assigned constant values across all steps (Fig. 6A), equivalent to the original stepping regulation model [1]. Likewise, during the steady-state periods before and after the lateral maneuver, all model parameters were also held constant to reflect steady-state walking [1].

As expected, allowing the stepping goals to adapt at each consecutive step emulated the experimentally observed stepping time series and errors, as well as the locations of the stepping distributions at each step of the lateral maneuver (Fig. 6B-D). However, these parameter variations did not influence the shape, area, or orientation of the stepping distributions, unlike what we observed experimentally (Fig. 6D-E). Thus, allowing the stepping goals to adapt from step to step appears necessary, but is not sufficient to elicit experimentally plausible stepping dynamics during lateral maneuvers.
Next, we added the ability to adaptively modulate $\rho$ at each step (Fig. 7A) to incorporate our theoretical predictions for an idealized four-step lane-change maneuver (Figs. 4C & 4F). Typical steady-state walking heavily weights regulating step width over lateral position (i.e., $\rho \approx 0.9$) [1], presumably to maintain lateral stability. Conversely, we expect people to trade off stability to gain maneuverability [8, 9] while executing this maneuver. We therefore set $\rho = 0.50$ at the transition step (Fig. 7A) to specify equal weighting of step
width and position regulation (Fig. 4F), thereby maximizing maneuverability. We then set $\rho = 0.7$ at the preparatory and recovery steps (Fig. 7A) to reflect an intermediate multi-objective cost weighting (Fig. 4F). The same adaptive stepping goals (Fig. 6) were again incorporated here. All other model parameters were assigned constant values across all steps (Fig. 7A).

Therefore, in addition to adapting both the stepping goals (Fig. 6) and $\rho$ (Fig. 7) from step to step, we then doubled the additive noise ($\sigma_a$) in the model at the preparatory and transition steps (Fig. 8A). Additive noise is
thought to reflect physiologic noise from a variety of sources, including sensory, perceptual, and/or motor processes [29]. Here, we assessed whether increasing this additive noise could emulate the increases in the stepping distribution areas observed experimentally (Fig. 5B-C).

Allowing the stepping goals, $\rho$, and $\sigma_a$ to adapt at each consecutive step again emulated the experimentally observed stepping time series, stepping errors, and stepping distribution locations at each step of the lateral maneuver (Fig. 8B-D) and qualitatively captured the shapes of the stepping distributions (Fig. 8D-E).
Increasing $\sigma_a$ also increased the areas of the stepping distributions at the preparatory, transition, and recovery steps (Fig. 8D-E). Interestingly, increasing $\sigma_a$ also influenced the orientations of these distributions, inducing a clockwise rotation at the preparatory step and a counterclockwise rotation at the transition step (Fig. 8D-E). These rotations reflect the individual steps at which the increased variability was applied. Increasing the noise at the preparatory step (shown here as taken with the left foot) can only increase variability in the horizontal direction in the $[z_L, z_R]$ plane. Conversely, increasing the noise at the subsequent transition step (taken with the right foot) can only increase variability in the vertical direction.

We designed these simulations specifically to emulate our theoretical predictions for an idealized four-step lane-change maneuver (Figs. 4C & 4F). Taken together, the results demonstrate that adapting the stepping goals ($z_B^*, w^*$; Fig. 6), the relative weight given to regulation for of those goals ($\rho$; Fig. 7), and the additive noise ($\sigma_a$; Fig. 8) allowed our simulations to emulate the key changes in stepping dynamics observed during this lateral maneuver task (Fig. 5).

**DISCUSSION**

Understanding how humans perform accurate, goal-directed walking movements in the face of inherent variability, redundancy, and equifinality remains a fundamental question in human motor neuroscience. In pursuit of this aim, we previously developed a theoretical framework to describe how humans regulate stepping to achieve continuous, straight-ahead walking [1]. Models developed from this framework provide goal-directed “stepping regulation templates” that are both analogous and complimentary to mechanical templates that describe the within-step mechanics and dynamics of walking (e.g., [22, 23, 32, 33]). Here, because humans rarely perform long bouts of steady-state walking [51, 52], we therefore examined how our basic regulation template scheme could be used to model human stepping dynamics during a prescribed lateral maneuver that, by necessity, strongly deviated from steady state motion.

Our key theoretical contribution is that we show how our previously developed theoretical stochastic optimal control framework can model how humans adapt their lateral stepping to enact non-steady-state lateral maneuvers (Fig. 4). Specifically, we show this can be accomplished by adding an additional layer to our previous control hierarchy: namely, the processes responsible for adapting, on a step-by-step basis, the cost landscapes governing stepping regulation. Our key empirical contribution is we then demonstrate (Fig. 5) that humans do indeed execute lateral maneuvers in a manner consistent with our theoretical predictions. Our key computational contribution is that we show that allowing the parameters in our lateral stepping regulation model to adapt from each step to the next can emulate the changes in lateral stepping dynamics exhibited by humans (Figs. 6-8). To our knowledge, our results are the first to demonstrate that humans use evolving cost landscapes in real time to perform such an adaptive motor task and, furthermore, that such adaptation can occur quickly – over only one step.
In our simulations (Figs. 6-8), we chose model parameters based on idealized theoretical considerations (Fig. 4), only loosely related to general trends observed in our experiment. While our resulting simulations did not precisely “fit” the experimental data, they were not intended to. On the contrary, our aim was to demonstrate that reasonable values of the parameters could yield the same basic dynamical and statistical stepping features that humans exhibited during this lateral maneuver task (Fig. 8). This approach is directly analogous to using mechanical “templates” of within-step dynamics of legged locomotion to reveal basic principles of walking and to propose fundamental hypotheses about what high-level control strategies might be acting [59, 60].

Here, we used our theoretical framework to derive such hypotheses (Fig. 4). We then used both experiments (Fig. 5) and simulations (Figs. 6-8) to validate the theoretical predictions facilitated by our templates [1] for an entirely new locomotor context.

Many studies have used optimal control theory to model a myriad of individual motor tasks (e.g., [61-65]). Such efforts, however, did not consider tasks where the task goals and/or movement objectives change as the task is being performed. Likewise, many studies have addressed motor adaptation (e.g., [66, 67]) and/or motor learning (e.g., [68, 69]), including for walking tasks (e.g., [70-72]). These paradigms, however, track how task performance changes slowly over many repetitions (typically at least 10’s to 100’s), and not from one repetition to the next. Furthermore, while multiple studies have demonstrated that such longer-term adaptation can be replicated by “lag-1” type computational models that explicitly correct only for errors experienced on one previous iteration of the task [73-79], those models were not themselves “adaptive”. On the contrary, they presumed some constant process (with constant model parameters) that achieved adaptation over multiple repetitions of the task under consideration. In sum, none of these extremely well-trodden paradigms quite capture the nature of the task we studied here.

Conversely, rapid adaptation in response to changing environmental contexts has been demonstrated in both birdsong [80-82] and human speech [83]. This apparent plasticity of a well learned, crystalized behavior suggests that vocalization may be controlled by a “malleable template,” in which trial-by-trial variability is used to adapt learned behaviors [84]. Such rapid adaptations have also been observed in both animal [15, 16, 85-88] and human [7-9] locomotion (i.e., maneuvers). As the associated motor planning processes occur nearly instantaneously [89-91], this prior work supports our findings that humans can and do make rapid adaptations to their stepping regulation to enact lateral maneuvers. Thus, in demonstrating that our lateral stepping regulation framework successfully predicts how humans perform lateral maneuvers, our findings support the notion that humans use a malleable template [84] to make “embodied decisions” [6] about how to regulate their stepping movements in real time during ongoing locomotion.

Prior work has postulated the existence of a “stability-maneuverability trade-off” during lateral maneuvers [7,
However, in the absence of a coherent, predictive, theoretical framework, this stability-maneuverability trade-off has not been adequately defined, much less confirmed. We propose that our lateral stepping regulation framework, and the models derived from it, provides the necessary theoretical and computation foundation needed to describe how humans trade-off stability for maneuverability during lateral movements. By extension, our findings suggest that stability and maneuverability in the context of locomotion are not distinct and independent concepts, but rather different manifestations of the same underlying stepping regulation process. Indeed, if stability and maneuverability were independent, as often assumed, humans should be able to remain stable and maneuverable simultaneously. However, our theoretical framework (Fig. 4) demonstrates precisely how and why humans must trade-off some degree of stability (i.e., $w$-regulation) to gain the maneuverability needed to perform lateral maneuvers.

Walking in the real world often requires maneuverability to adapt to changing environmental conditions or goals. We suggest that such maneuvers are governed by a hierarchical control/regulation schema with at least three distinct layers: low-level processes that govern within-step dynamics to ensure viability [34, 59, 60], step-to-step regulation to achieve goal-directed walking [1, 29, 34, 36], and the presently demonstrated modulation of stepping regulation to achieve adaptability. The ability of our lateral stepping regulation framework to emulate human stepping during the lane change maneuver studied here demonstrates that its predictive capabilities extend to a much greater range of walking tasks than initially thought, encompassing not just steady state walking, but transient behaviors as well.

METHODS

Ethics Statement
Prior to participating, all participants signed informed consent statements approved by the Institutional Review Board at The Pennsylvania State University.

Lateral Maneuvers in Humans
The data analyzed here were collected as part of a previous experiment involving twenty young healthy adults (Table 1) [58]. All participants were screened to ensure they had no history of orthopedic problems, recent lower extremity injuries, any visible gait anomalies, or were taking medications that may have influenced their gait.

Table 1 – Participant Characteristics. All values except Sex are given as Mean ± Standard Deviation.

| Characteristic: | Value:          |
|----------------|----------------|
| Sex            | 9 M / 11 F     |
| Age [yrs]      | 21.7 ± 2.6     |
Body Height [m] \hspace{1cm} 1.73 \pm 0.08
Body Mass [kg] \hspace{1cm} 69.9 \pm 12.1
Body Mass Index (BMI) [kg/m²] \hspace{1cm} 23.2 \pm 2.7
Leg Length [m] \hspace{1cm} 0.82 \pm 0.06

The experimental protocols were described in detail previously [58]. Briefly, participants walked in an “M-Gait” system, comprised of a 1.2m wide motorized treadmill in a virtual reality environment (Motek, Amsterdam, Netherlands). Each participant walked at a constant speed of 0.75 m/s. Following a 4-minute acclimation trial, participants completed several different walking trials involving path navigation. The data analyzed here were generated from one such trial, during which participants were instructed to switch between two parallel paths, centered 0.6m apart, following an audible cue (Fig. 2). Participants completed 6 maneuvers during one 4-minute walking trial, and were instructed to walk normally on their current path between maneuvers. The first and last maneuvers occurred too close to the beginning and end of the walking trial, respectively, to ensure participants were walking normally both before and after the maneuver.

Therefore, the present analysis included only the middle four maneuvers from each participant, for a total of 80 lateral maneuvers.

Kinematic data were recorded from 16 retroreflective markers placed on the head, pelvis, and feet of each participant. Marker trajectories were collected at 120Hz using a 10-camera Vicon motion capture system (Oxford Metrics, Oxford, UK) and post-processed using Vicon Nexus and D-Flow software (Motek, Amsterdam, Netherlands). Marker trajectories were analyzed in MATLAB (MathWorks, Natick, MA). Heel strikes were determined using a velocity-based detection algorithm [92]. Lateral foot placements ($z_L$ and $z_R$) were defined as the lateral location of the heel marker at each step. Step width ($w$) and lateral position ($z_B$) were then determined at each step using Eq (1) (Fig. 1C).

We were unable to determine accurate heel strikes for one lateral maneuver and consequently excluded this maneuver from further analyses. Stepping data from the remaining 79 maneuvers were normalized to a constant direction (left-to-right) and to the initiation of the transition, defined here as the last step taken on the original path. Means and standard deviations of step width and position were determined at each step of the lateral maneuver. The steady-state mean step width was determined from all participants and all steps, excluding the 5 steps before to 10 steps after each transition. The steady-state mean position was determined similarly, but was determined separately for each path (left and right). These steady-state means were used to define the stepping goals for this task, [$z_B^*$, $w^*$]: $w^*$ was defined as the steady-state mean step width, and $z_B^*$ before and after the maneuver was defined as the steady-state mean position on the left path and the steady-state mean position on the right path, respectively. Errors in both position and step width (mean±SD) were determined at each step of each maneuver as the differences in each from these steady-state stepping goals (Fig. 2C).
Model of Lateral Stepping Regulation

To model lateral stepping, the simplest, mechanically sufficient biped includes the lateral locations of the center-of-mass and each of the two feet ($z_{Ln}$ and $z_{Rn}$; Fig. 1B) [32, 93, 94]. We presume that left and right foot placement are coordinated to achieve some more general walking task goal or goals. One such goal is to maintain lateral balance by regulating step width ($w_n$; Fig. 1C) [23, 95-97]. Humans also regulate the lateral position of the body’s mass center with respect to their path [18, 94, 98, 99], approximated by the midpoint between the two feet ($z_{Bn}$; Fig. 1C) during upright walking [93, 95]. By selecting appropriate foot placements, $\{z_L, z_R\}$, humans can regulate both their center of mass position and step width, $\{z_{Bn}, w_n\}$ [1], via:

$$
\begin{bmatrix}
  z_{Bn} \\
  w_n
\end{bmatrix} =
\begin{bmatrix}
  \frac{1}{2} & \frac{1}{2} \\
  -1 & 1
\end{bmatrix}
\begin{bmatrix}
  z_{Ln} \\
  z_{Rn}
\end{bmatrix}
$$  

(1)

During continuous straight-ahead walking, multi-objective regulation of primarily step width and secondarily lateral position captures human lateral step-to-step dynamics [1]. The regulation model minimizes errors with respect to both step width and lateral position using the goal functions [37, 38, 40]:

$$
F_{zB} = z_{Bn} - z_B^*
$$

$$
F_w = w_n - w^*
$$  

(2)

where the goal is to drive each $F \rightarrow 0$. The value of each regulated variable at the subsequent step is determined independently using the state update equations:

$$
z_{B(n+1)} = z_{Bn} + g \left(1 + \sigma_{zm} \right) u_{zB} \left(z_{Bn} \right) + \sigma_{zB} v_{zB}
$$

$$
w_{n+1} = w_n + g \left(1 + \sigma_{wm} \right) u_{w} \left(w_n \right) + \sigma_{wa} v_{wa}
$$  

(3)

where $\sigma_{zm}$ and $\sigma_{wm}$ represent multiplicative and additive noise terms, respectively.

The control inputs, $u_{zB}(z_{Bn})$ and $u_{w}(w_n)$, were derived analytically as stochastic optimal single-step regulators with direct error feedback [1, 29, 30, 40], following the Minimum Intervention Principle [42, 64]. Such controllers are optimal with respect to the following quadratic cost functions:

$$
C_{zB} = \alpha e_{zB(n+1)}^2 + \gamma u_{zB}^2
$$

$$
C_w = \alpha e_{w(n+1)}^2 + \gamma u_{w}^2
$$  

(4)
The first term of each cost function penalizes errors with respect to the goal function (Eq (2)) at the next step. The second term penalizes “effort”, quantified as the magnitude of the control input. Here, $\alpha$ and $\gamma$ were positive constants that weighted the two terms of each cost function [1]. The subsequent analytical derivation [1] yielded control inputs as:

$$u_{zB}(z_{Bn}) = -\frac{1}{1 + \sigma_m^2 + (\gamma/\alpha)} \left(z_{Bn} - z^*_B\right)$$

$$u_w(w_n) = -\frac{1}{1 + \sigma_m^2 + (\gamma/\alpha)} \left(w_n - w^*\right)$$

The value of each regulated variable, $[z_B, w]$, at the subsequent step is determined independently using separate, individual optimal controllers. Thus, two corresponding foot placements are determined, one using the predicted value of position, and the other using the predicted value of step width. The final predicted value of foot placement is determined as a weighted average ($\rho$) of the predictions from each regulated variable, where $\rho = 0$ indicates 100% position control and $\rho = 1$ indicates 100% step width control [1]. Hence, stepping regulation is conceived here as arising from a “mixture of experts” [63, 100]. In this way, the value of $\rho$ directly determines the shape of the cost landscape, as evidenced by the changing shapes of the variability ellipses in the $[z_L, z_R]$ plane (Fig. 7).

As in our prior work [1], the following parameters were set to constant values for all simulations. The baseline amplitude of the additive noise ($\sigma_{-\alpha}$) was set to the steady-state standard deviation of each variable. The amplitude of the multiplicative noise ($\sigma_m$) was set to 10% of the additive noise level (i.e., $0.1 \cdot \sigma_{-\alpha}$). The magnitude of the additive and multiplicative noise at a given step were determined from the Gaussian random variables $\nu_{-\alpha}$ and $\nu_m$, respectively, each with zero mean and unit variance. Error correction was weighted much more heavily than effort in the cost function by setting $\gamma/\alpha = 0.10$.

**Steady-State Stepping Regulation**

We determined if any constant value of $\rho$ could emulate human stepping throughout the lateral maneuver (Fig. 3C). For each value of $\rho$ between $\rho = 0.00$ and $\rho = 1.00$, in increments of $\Delta \rho = 0.01$, we conducted 1000 batches of 1000 simulations of the lateral maneuver. For each batch, we determined the means and standard deviations of both position and step width at each step. We then computed the overall mean and standard deviation of each variable at each step across all batches. For each variable, simulated and experimental values were compared by plotting the simulated overall mean $\pm$ 1SD against 95% confidence intervals of the experimental data determined from bootstrapping [40, 101]. For a given value of $\rho$, we considered the model
successful at capturing human stepping if the simulated means and standard deviations of step width and position fell within the corresponding experimental 95% confidence intervals at every step of the lateral maneuver (Fig. 3C).

Stepping Regulation for Non-Steady-State Tasks

When the $z_B^*$ and $w^*$ GEMs are viewed in the $[z_L, z_R]$ plane, it is clear that at least one intermediate step is necessary to accomplish any $\Delta z_B^*$ and/or $\Delta w^*$ maneuver (Fig. 4A). For the minimum two-step maneuver strategy (Fig. 4B), the stepping goals for either transition step can be determined algebraically from the steady-state initial, $[z_{Bi}^*, w_{i}^*]$, and final, $[z_{Bi}^*, w_{f}^*]$, stepping goals. We first determined the initial and final foot placement goals from the initial and final stepping goals using Eq (1). The foot used to take the intermediate step must be placed at that foot’s final foot placement goal, while the stance foot remains at its initial foot placement goal. For example, if the right foot is used to take the intermediate step (Fig. 4B; a), the intermediate foot placement goals are $[z_{Li}^*, z_{Rf}^*]$. By transforming these intermediate foot placement goals back into $[z_B, w]$ coordinates using Eq (1), and re-defining the final stepping goals as the initial stepping goals plus the changes in the stepping goals, $[\Delta z_B^*, \Delta w^*]$, the intermediate stepping goals are:

$$z_B^* = \frac{1}{2}(z_{Li}^* + z_{Rf}^*) = \frac{1}{2}(z_{Bi}^* - \frac{1}{2} w_{i}^* + \frac{1}{2}(z_{Bi}^* + \Delta z_B^*) + \frac{1}{2}(w_{i}^* + \Delta w^*))$$

$$w^* = (z_{Rf}^* - z_{Li}^*) = (z_{Bi}^* + \Delta z_B^*) + \frac{1}{2}(w_{i}^* + \Delta w^*) - (z_{Bi}^* - \frac{1}{2} w_{i}^*)$$

The same procedure can be used to determine the stepping goals for an intermediate step taken with the left foot (Fig. 4B; b) by simply reversing the initial and final components of the intermediate foot placement goals and making the corresponding substitutions in Eq (6).

However, equifinality exists in the number of steps and their placements that can be used to accomplish any $\Delta z_B^*$ and/or $\Delta w^*$ maneuver. Experimentally, we most commonly observed a four-step maneuver strategy with distinct preparatory, transition, and recovery steps (Figs. 2, 4C). To determine the stepping goals associated with an idealized version of such a four-step strategy, we first calculated a single intermediate offset (ε) by averaging the experimental differences in foot placement at each preparatory and recovery step relative to steady-state walking. Assuming a left-to-right transition with preparatory and recovery steps that were each taken with the left foot, the foot placement goals at the preparatory, $[z_{Li}^{*pr}, z_{Ri}^{*pr}]$, and recovery, $[z_{Li}^{*rc}, z_{Ri}^{*rc}]$, steps were defined as $[z_{Li}^* + \epsilon, z_{Ri}^*]$ and $[z_{Li}^*, z_{Ri}^* - \epsilon]$, respectively. These foot placement goals were then transformed back into $[z_B, w]$ coordinates using Eq (1) to determine the preparatory, $[z_B^{*pr}, w^{*pr}]$, and...
recovery, \([z^{*}_{B,Re}, w^{*}_{Re}]\), stepping goals. The transition stepping goals, \([z^{*}_{B,Tr}, w^{*}_{Tr}]\), were then determined using

Eq (6) above, substituting the preparatory and recovery stepping goals for the initial and final stepping goals.

The stepping goals themselves, however, do not provide insight into how humans regulated their stepping
during the lateral maneuver. For a four-step maneuver strategy with a large primary transition step and
smaller preparatory and recovery steps, as was observed experimentally (Fig. 2), stepping distributions are
theoretically predicted to be most isotropic at the primary transition step and intermediately isotropic at the
preparatory and recovery steps.

We qualitatively assessed the accuracy of the theoretically predicted stepping goals and distributions by
comparing them to the experimental data at each step, plotted in the \([z_{L}, z_{R}]\) plane (Fig. 5B). The maneuver
completed with a large cross-over step was excluded from analyses of the stepping distributions, as this
maneuver was completed with a different stepping strategy. Using the remaining 78 maneuvers, we first
computed the covariance matrix of left and right foot placements at each step: \(C = \text{cov}(z_{L}, z_{R})\). We take \(\{\lambda_1, \lambda_2\}\) and \(\{e_1, e_2\}\) to denote the eigenvalues and eigenvectors of \(C\), respectively. We then construct a 95% prediction ellipse by scaling the eigenvalues by the 95\(^{th}\) percentile critical value of the Chi-Squared distribution:

\[
\begin{bmatrix}
z_L(t) \\
z_R(t)
\end{bmatrix} =
\begin{bmatrix}
e_1 & e_2
\end{bmatrix}
\begin{bmatrix}
\sqrt{5.991\lambda_1}\cos(t) \\
\sqrt{5.991\lambda_2}\sin(t)
\end{bmatrix}, \quad \text{where: } t = 0...2\pi.
\]

Thus, we expect 95% of the data points to lie inside this ellipse, assuming a bivariate normal distribution. We
then characterized each such ellipse by its shape, size, and orientation (Fig. 5C). We defined the shapes of
each ellipse by their aspect ratio, computed as the ratio of the major- and minor-axis eigenvalues: \(\lambda_1/\lambda_2\). We
computed the sizes of each ellipse as their area:

\[
A = 5.991\pi\sqrt{\lambda_1\lambda_2}.
\]

We computed the orientations of each ellipse as the angle of the major axis measured counterclockwise from
the \(w^*\) GEM:
Adaptive Stepping Regulation for Lateral Maneuvers

We adapted parameters of our goal-directed multi-objective lateral stepping regulation model from step-to-step to determine if such adaptions could model the observed stepping dynamics during the lateral maneuver task. For each model iteration, all parameters not explicitly modulated were assigned a constant value in accordance with our previous steady-state stepping regulation model [1]. We first replaced the stepping goals $z^*_{B}$ and $w^*$ in the model (Eq (2)) with the estimated adaptive stepping goals (see previous section) (Fig. 6A).

We next adapted $\rho$, the model parameter that specifies the relative weighting of step width and position regulation. The values assigned to $\rho$ were chosen based upon theoretical predictions of the maneuverability and error correction requirements at each intermediate step (see previous section; Fig. 5). Specifically, we set $\rho = 0.50$ at the transition step, specifying equal weighting of step width and position regulation. We set $\rho = 0.7$ at the preparatory and recovery steps, a value intermediate between the approximate value observed for steady-state walking ($\rho \approx 0.9$) and the value of 0.5 used for the transition step (Fig. 7A).

Finally, in addition to the adaptive stepping goals and $\rho$ modulation, we doubled the additive noise parameter ($\sigma_a$) at the preparatory and transition steps (Fig. 8A). The increased area of the stepping distributions at the preparatory, transition, and recovery steps reflects greater overall variability with respect to both the $w^*$ and $z^*_{B}$ GEMs. Additive noise is thought to reflect physiologic noise from sensory, perceptual, and/or motor processes [29], which is expected to increase during the lateral maneuver task. Here, we increased the additive noise (Fig. 8A) to determine the extent to which this would qualitatively capture the types of increases in stepping distribution areas that we observed experimentally.

For each model iteration, we assessed the ability of the model to emulate the key changes in stepping dynamics observed experimentally: stepping time series (Figs. 6-8B), stepping errors (Figs. 6-8C), and stepping distributions (Figs. 6-8D-E) at each step of the lateral maneuver. The lateral maneuver was simulated 1000 times for each model iteration. All maneuvers were oriented from left-to-right, and the transition step was specified to be taken with the ipsilateral (i.e., right) foot relative to the direction of the transition.

\[
\Delta \theta = \begin{cases} 
\frac{\pi}{4} & \text{if } C_{2,1} = 0 \text{ and } C_{1,1} \geq C_{2,2} \\
\frac{3\pi}{4} & \text{if } C_{2,1} = 0 \text{ and } C_{1,1} < C_{2,2}, \\
\tan^{-1}\left(\lambda_1 - C_{1,1}, C_{2,1}\right) + \frac{\pi}{4} & \text{else}
\end{cases}
\]

where the $C_{ij}$ are the respective elements of the covariance matrix. For each ellipse characteristic computed at each step, we then calculated 95% confidence intervals for these values using bootstrapping.
consistent with all but one of the experimentally observed maneuvers. Time series (mean±SD) of foot placement ($z_L$ and $z_R$), position ($z_B$), and step width ($w$) were calculated at each step of the simulated maneuvers. Errors with respect to both position and step width (mean±SD) were calculated as the difference in the simulated position and step width relative to the stepping goals at each step of the simulated maneuvers. The simulated stepping time series and errors were compared to the middle 90% range of the experimental data at each step. The simulated stepping distributions were characterized by the aspect ratio, area, and orientation of a fitted 95% prediction ellipse (see previous section). Error bars at each step for each variable were calculated as 95% confidence intervals using bootstrapping. The simulated ellipse characteristics were compared to the bootstrapped 95% confidence intervals from the experimental data at each step for each variable.

**Statistical Comparisons**

As the experimental and simulated data were structured in very different ways, standard inferential statistics (e.g., t-test, ANOVA, etc.) would not be appropriate to compare these results. Furthermore, we could generate a sufficiently large number of model simulations to ensure small p-values for almost any comparison, thereby diminishing the comparative power of any such assessments. Instead, we used descriptive statistics (e.g., standard deviations, confidence intervals, etc.) to quantify the experimental observation values. We then compared model predictions to these observations. We inferred that model predictions that fell within experimentally observed standard deviations/confidence intervals were statistically consistent with the experimental results. Additionally, the aim of this analysis was to determine if hierarchical adaption of our goal-directed multi-objective lateral stepping regulation models could qualitatively emulate the same types of changes in stepping dynamics observed during the lateral maneuver task. Descriptive statistics were sufficient to accomplish this aim.
AUTHOR CONTRIBUTIONS

| Role                                | Authors         |
|-------------------------------------|-----------------|
| Conceptualization                   | DMD; JPC; JBD   |
| Data Curation                       | DMD; JBD        |
| Formal Analysis                     | DMD; JPC; JBD   |
| Funding Acquisition                 | JPC; JBD        |
| Investigation                       | DMD             |
| Methodology                         | DMD; JPC; JBD   |
| Project Administration              | JBD; JPC        |
| Resources                           | JPC; JBD        |
| Software                            | DMD; JPC; JBD   |
| Supervision                         | JPC; JBD        |
| Validation                          | DMD; JPC; JBD   |
| Visualization                       | DMD             |
| Writing – Original Draft Preparation| DMD             |
| Writing – Review & Editing          | DMD; JPC; JBD   |

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REFERENCES

1. Dingwell JB, Cusumano JP. Humans Use Multi-Objective Control to Regulate Lateral Foot Placement When Walking. PLoS Comput Biol. 2019;15(3):e1006850. doi: https://doi.org/10.1371/journal.pcbi.1006850.

2. Maeda RS, O'Connor SM, Donelan JM, Marigold DS. Foot placement relies on state estimation during visually guided walking. J Neurophysiol. 2017;117(2):480-91. doi: https://doi.org/10.1152/jn.00015.2016.

3. Moussaïd M, Helbing D, Garnier S, Johansson A, Combe M, Theraulaz G. Experimental study of the behavioural mechanisms underlying self-organization in human crowds. Proc Roy Soc B Biol Sci. 2009;276(1668):2755-62. doi: http://dx.doi.org/10.1098/rspb.2009.0405.

4. Matthys JS, Barton SL, Fajen BR. The critical phase for visual control of human walking over complex terrain. Proc Natl Acad Sci USA. 2017;114(32):E6720-E9. doi: https://doi.org/10.1073/pnas.1611699114.

5. Matthys JS, Yates JL, Hayhoe MM. Gaze and the Control of Foot Placement When Walking in Natural Terrain. Curr Biol. 2018;28(8):1224-33.e5. doi: https://doi.org/10.1016/j.cub.2018.03.008.

6. Gordon J, Maselli A, Lancia GL, Thiery T, Cisek P, Pezzulo G. The road towards understanding embodied decisions. Neuroscience & Biobehavioral Reviews. 2021;131:722-36. doi: https://doi.org/10.1016/j.neubiorev.2021.09.034.

7. Wu M, Matsubara JH, Gordon KE. General and Specific Strategies Used to Facilitate Locomotor Maneuvers. PLoS ONE. 2015;10(7):e0132707. doi: http://dx.doi.org/10.1371/journal.pone.0132707.

8. Acasio J, Wu MM, Fey NP, Gordon KE. Stability-maneuverability trade-offs during lateral steps. Gait Posture. 2017;52:171-7. doi: http://dx.doi.org/10.1016/j.gaitpost.2016.11.034.

9. Hsieh KL, Sheehan RC, Wilken JM, Dingwell JB. Healthy individuals are more maneuverable when walking slower while navigating a virtual obstacle course. Gait Posture. 2018;61:466-72. doi:
10. Barton SL, Matthys JS, Fajen BR. Visual regulation of gait: Zeroing in on a solution to the complex terrain problem. J Exp Psychol Hum Percep Perf. 2017;43(10):1773-90. doi: http://dx.doi.org/10.1037/xhp0000435. PubMed PMID: 1947580833; 2017-42972-005.

11. Kowalsky DB, Rebula JR, Ojeda LV, Adamczyk PG, Kuo AD. Human walking in the real world: Interactions between terrain type, gait parameters, and energy expenditure. PLoS ONE. 2021;16(1):e0228682. Epub 2021/01/14. doi: 10.1371/journal.pone.0228682. PubMed PMID: 33439858; PubMed Central PMCID: PMCPMC7806134.

12. Twardzik E, Duchowny K, Gallagher A, Alexander N, Strasburg D, Colabianchi N, et al. What features of the built environment matter most for mobility? Using wearable sensors to capture real-time outdoor environment demand on gait performance. Gait Posture. 2019;68:437-42.

13. Kazanski ME, Dingwell JB, Cusumano JP. How healthy older adults regulate lateral foot placement while walking in laterally destabilizing environments. J Biomech. 2020;104:109714.

14. Wu MM, Brown G, Gordon KE. Control of Locomotor Stability in Stabilizing and Destabilizing Environments. Gait Posture. 2017;55(Supplement C):191-8. doi: https://doi.org/10.1016/j.gaitpost.2018.02.015.

15. Birn-Jeffery AV, Hubicki CM, Blum Y, Renjewski D, Hurst JW, Daley MA. Don't break a leg: running birds from quail to ostrich prioritise leg safety and economy on uneven terrain. J Exp Biol. 2014;217(Pt 21):3786-96. Epub 2014/10/31. doi: 10.1242/jeb.102640. PubMed PMID: 25355848; PubMed Central PMCID: PMCPMC4213177.

16. Biewener AA, Bomphrey RJ, Daley MA, Ijspeert AJ. Stability and manoeuvrability in animal movement: lessons from biology, modelling and robotics. Proc Roy Soc B Biol Sci. 2022;104:2286-42. doi:doi:10.1098/rspb.2021.2492.

17. Othayoth R, Xuan Q, Wang Y, Li C. Locomotor transitions in the potential energy landscape-dominated regime. Proc Roy Soc B Biol Sci. 2021;288(1949):20202734. doi:10.1098/rspb.2020.2734.

18. Bruijn SM, van Dieën JH. Control of human gait stability through foot placement. J R Soc Interface. 2018;15(143):1-11. doi: https://doi.org/10.1098/rsif.2017.0816.

19. Bauby CE, Kuo AD. Active Control of Lateral Balance in Human Walking. J Biomech. 2000;33(11):1433-40. doi: http://dx.doi.org/10.1016/S0021-9290(00)00101-9.

20. MacKinnon CD, Winter DA. Control of Whole Body Balance In The Frontal Plane During Human Walking. J Biomech. 1999;26(6):633-44. doi: https://doi.org/10.1016/0021-9290(99)0027-C.

21. Townsend MA. Biped Gait Stabilization Via Foot Placement. J Biomech. 1985;18(1):21-38. doi: http://dx.doi.org/10.1016/0021-9290(85)90042-9.

22. Hof AL. The 'extrapolated center of mass' concept suggests a simple control of balance in walking. Hum Mov Sci. 2008;27(1):112-25. doi: https://doi.org/10.1016/j.humov.2007.08.003.

23. Kuo AD. Stabilization of Lateral Motion in Passive Dynamic Walking. Int J Robotics Res. 1999;18(9):917-30. doi: https://doi.org/10.1177/0278364992066655.

24. Kelsey JL, Browner WS, Seeley DG, Nevitt MC, Cummings SR. Risk factors for fractures of the distal forearm and proximal humerus. The Study of Osteoporotic Fractures Research Group. Am J Epidemiol. 1992;135(5):477-89. Epub 1992/03/01. doi: 10.1093/oxfordjournals.aje.a116314. PubMed PMID: 1570814.

25. Parkkari J, Kannus P, Palvanen M, Natri A, Vainio I, Aho H, et al. Majority of Hip Fractures Occur as a Result of a Fall and Impact on the Greater Trochanter of the Femur: A Prospective Controlled Hip Fracture Study with 206 Consecutive Patients. Calcif Tissue Int. 1999;65(3):183-7. doi: http://dx.doi.org/10.1007/s0022399000679.

26. Winter DA. Kinematic and Kinetic Patterns in Human Gait: Variability and Compensating Effects. Hum Mov Sci. 1984;3(1-2):51-76. doi: http://dx.doi.org/10.1016/0167-9457(84)90005-8.

27. Kang HG, Dingwell JB. Separating the Effects of Age and Speed on Gait Variability During Treadmill Walking. Gait Posture. 2008;27(4):572-7. doi: https://doi.org/10.1016/j.gaitpost.2007.07.009.

28. Bernstein N. The Coordination and Regulation of Movements. New York, NY: Pergamon Press; 1967.

29. Dingwell JB, John J, Cusumano JP. Do Humans Optimally Exploit Redundancy to Control Step Variability in Walking? PLoS Comput Biol. 2010;6(7):e1000856. doi: http://dx.doi.org/10.1371/journal.pcbi.1000856.
31. Dingwell JB, Salinas MM, Cusumano JP. Increased Gait Variability May Not Imply Impaired Stride-To-Stride Control of Walking in Healthy Older Adults. Gait Posture. 2017;55:131-7. doi: http://dx.doi.org/10.1016/j.gaitpost.2017.03.018.

32. Hobbelten DGE, Wisse M. Active Lateral Foot Placement for 3D Stabilization of a Limit Cycle Walker Prototype. Int J Hum Robot. 2009;6(1):93-116. doi: http://dx.doi.org/10.1142/S0219843609001632.

33. Koolen T, de Boer T, Rebula J, Goswami A, Pratt J. Capturability-based analysis and control of legged locomotion, Part 1: Theory and application to three simple gait models. Int J Robotics Res. 2012;31(9):1094-113. doi: https://doi.org/10.1177/0278364912452673.

34. Patil NS, Dingwell JB, Cusumano JP. Viability, task switching, and fall avoidance of the simplest dynamic walker. bioRxiv. 2022:2022.01.16.476517. doi: 10.1101/2022.01.16.476517.

35. Zaytsev P, Wolfslag W, Ruina A. The Boundaries of Walking Stability: Viability and Controllability of Simple Models. IEEE Trans Robot. 2018;34(2):336-52. doi: 10.1109/TRO.2017.2782818.

36. Patil NS, Dingwell JB, Cusumano JP. Task-level regulation enhances global stability of the simplest dynamic walker. J R Soc Interface. 2020;17(168):20200278. Epub 2020/07/18. doi: 10.1098/rsif.2020.0278. PubMed PMID: 32674710; PubMed Central PMCID: PMCPMC7423424.

37. Cusumano JP, Cesari P. Body-Goal Variability Mapping in an Aiming Task. Biol Cybern. 2006;94(5):367-79. doi: https://doi.org/10.1007/s00422-006-0052-1.

38. Cusumano JP, Dingwell JB. Movement Variability Near Goal Equivalent Manifolds: Fluctuations, Control, and Model-Based Analysis. Hum Mov Sci. 2013;32(5):899-923. doi: http://dx.doi.org/10.1016/j.humov.2013.07.019.

39. Cusumano JP, Mahoney JM, Dingwell JB. The Dynamical Analysis of Inter-Trial Fluctuations Near Goal Equivalent Manifolds. Adv Exp Med Biol. 2014;826:125-45. doi: http://dx.doi.org/10.1007/978-1-4939-1338-1_9.

40. John J, Dingwell JB, Cusumano JP. Error Correction and the Structure of Inter-Trial Fluctuations in a Redundant Movement Task. PLoS Comput Biol. 2016;12(9):e1005118. doi: http://dx.doi.org/10.1371/journal.pcbi.1005118.

41. Todorov E. Optimality principles in sensorimotor control. Nat Neurosci. 2004;7(9):907-15. doi: http://dx.doi.org/10.1038/nn1309.

42. Todorov E, Jordan MI. Optimal feedback control as a theory of motor coordination. Nat Neurosci. 2002;5(11):1226-35. doi: https://doi.org/10.1038/nn963.

43. Render AC, Kazanski ME, Cusumano JP, Dingwell JB. Walking humans trade off different task goals to regulate lateral stepping. J Biomech. 2021;119:110314. Epub 2021/03/06. doi: 10.1016/j.jbiomech.2021.110314. PubMed PMID: 33667882; PubMed Central PMCID: PMCPMC8081051.

44. Bohnsack-McLagan NK, Cusumano JP, Dingwell JB. Adaptability of Stride-To-Stride Control of Stepping Movements in Human Walking. J Biomech. 2016;49(2):229-37. doi: https://doi.org/10.1016/j.jbiomech.2015.12.010.

45. Dingwell JB, Bohnsack-McLagan NK, Cusumano JP. Humans Control Stride-To-Stride Stepping Movements Differently for Walking and Running, Independent of Speed. J Biomech. 2018;76:144-51. doi: https://doi.org/10.1016/j.jbiomech.2018.05.034.

46. Decker LM, Cignetti F, Potter JF, Studenski SA, Stergiou N. Use of Motor Abundance in Young and Older Adults during Dual-Task Treadmill Walking. PLoS ONE. 2012;7(7):e41306. doi: http://dx.doi.org/10.1371/journal.pone.0041306.

47. Terrier P, Dérijaz O. Persistent and anti-persistent pattern in stride-to-stride variability of treadmill walking: Influence of rhythmic auditory cueing. Hum Mov Sci. 2012;31(6):1585-97. doi: http://dx.doi.org/10.1016/j.humov.2012.05.004.

48. Roerdink M, Daffertshofer A, Marmelat V, Beek PJ. How to Sync to the Beat of a Persistent Fractal Metronome without Falling Off the Treadmill? PLoS ONE. 2015;10(7):e0134148. doi: http://dx.doi.org/10.1371/journal.pone.0134148.

49. Terrier P. Fractal Fluctuations in Human Walking: Comparison Between Auditory and Visually Guided Stepping. Ann Biomed Eng. 2016;44(9):2785-93. doi: http://dx.doi.org/10.1007/s10439-016-1573-y.

50. Roerdink M, de Jonge CP, Smidt LM, Daffertshofer A. Tightening Up the Control of Treadmill Walking: Effects of
51. Orendurff MS, Schoen JA, Bernatz GC, Segal AD, Klute GK. How Humans Walk: Bout Duration, Steps per Bout, and Rest Duration. J Rehabil Res Develop. 2008;45(7):1077-90. doi: http://dx.doi.org/10.1682/JRRD.2007.11.0197.

52. Lugade V, Kuntapun J, Prupetkaew P, Boripuntakul S, Verner E, Silsupadol P. Three-Day Remote Monitoring of Gait Among Young and Older Adults Using Participants’ Personal Smartphones. J Aging Phys Act. 2021;29(6):1026-33. Epub 2021/08/05. doi: 10.1123/japa.2020-0353. PubMed PMID: 34348231.

53. Robinovitch SN, Feldman F, Yang Y, Schonnop R, Leung PM, Sarraf T, et al. Video capture of the circumstances of falls in elderly people residing in long-term care: an observational study. Lancet. 2013;381(9860):47-54. doi: http://dx.doi.org/10.1016/S0140-6736(12)61263-X.

54. Matthies JS, Fajen BR. Visual control of foot placement when walking over complex terrain. J Exp Psychol Hum Percep Perf. 2014;40(1):106-15. doi: http://psycnet.apa.org/doi/10.1037/a0033101.

55. Marigold DS, Patla AE. Gaze fixation patterns for negotiating complex ground terrain. Neuroscience. 2007;144(1):302-13. doi: https://doi.org/10.1016/j.neuroscience.2006.09.006.

56. Land MF. Eye movements and the control of actions in everyday life. Prog Retin Eye Res. 2006;25(3):296-324. Epub 2006/03/07. doi: 10.1016/j.preteyeres.2006.01.002. PubMed PMID: 16516530.

57. Patla AE, Vickers JN. How far ahead do we look when required to step on specific locations in the travel path during locomotion? Exp Brain Res. 2003;148(1):133-8. doi: https://doi.org/10.1007/s00221-002-1246-y.

58. Kazanski ME, Dingwell JB. Effects of age, physical and self-perceived balance abilities on lateral stepping adjustments during lateral balance tasks. Gait Posture. 2021;88:311-7. Epub 2021/06/26. doi: 10.1016/j.gaitpost.2021.05.025. PubMed PMID: 34171786.

59. Dickinson MH, Farley CT, Koehl MA, Kram R, Lehman S. How Animals Move: An Integrative View. Science. 2000;Apr 7; 288(5463):100-6. doi: https://doi.org/10.1126/science.288.5463.100.

60. Full RJ, Koditschek DE. Templates and Anchors: Neuromechanical Hypothesis of Legged Locomotion on Land. J Exp Biol. 1999;202(23):3325-32.

61. Engelbrecht SE. Minimum Principles in Motor Control. J Math Psychol. 2001;45(3):497-542. doi: https://doi.org/10.1006/jmps.2000.1295.

62. Collins JJ. The Redundant Nature of Locomotor Optimization Laws. J Biomech. 1995;28(3):251-67. doi: https://doi.org/10.1016/0021-9290(94)00072-C.

63. Wolpert DM, Ghahramani Z. Computational Principles of Movement Neuroscience. Nat Neurosci. 2000;3(Supp):1212-7. doi: 10.1038/81497.

64. Scott SH. Optimal Feedback Control and the Neural Basis of Volitional Motor Control. Nat Rev Neurosci. 2004;5(7):532-46. doi: https://doi.org/10.1038/nrn1427.

65. Guigon E. A computational theory for the production of limb movements. Psychol Rev. 2021:No Pagination Specified-No Pagination Specified. doi: 10.1037/rev0000323.

66. Wolpert DM, Diedrichsen J, Flanagan JR. Principles of sensorimotor learning. Nat Rev Neurosci. 2011;12(12):739-51. doi: 10.1038/nrn3112.

67. Shadmehr R, Smith MA, Krakauer JW. Error Correction, Sensory Prediction, and Adaptation in Motor Control. Annual Review of Neuroscience. 2010;33(1):89-108. doi: http://dx.doi.org/10.1146/annurev-neuro-060909-153135.

68. Haith AM, Krakauer JW. The multiple effects of practice: skill, habit and reduced cognitive load. Curr Opin Behav Sci. 2018;20:196-201. Epub 2019/04/05. doi: 10.1016/j.cobeha.2018.01.015. PubMed PMID: 30944847; PubMed Central PMCID: PMCPMC6443249.

69. Krakauer JW, Hadjiosif AM, Xu J, Wong AL, Haith AM. Motor Learning. Compr Physiol. 2019;9(2):613-63. Epub 2019/03/16. doi: 10.1002/cphy.c170043. PubMed PMID: 30873583.

70. Choi JT, Bastian AJ. Adaptation reveals independent control networks for human walking. Nat Neurosci. 2007;10(8):1055-62. doi: https://doi.org/10.1038/nn1930.

71. Reisman DS, Block HJ, Bastian AJ. Interlimb Coordination During Locomotion: What Can be Adapted and Stored?
72. Abram S, Selinger J, Donelan J. Energy optimization is a major objective in the real-time control of step width in human walking. J Biomech. 2019;85:91.

73. Scheidt RA, Dingwell JB, Mussa-Ivaldi FA. Learning to Move Amid Uncertainty. J Neurophysiol. 2001;86(2):971-85.

74. van Beers RJ, van der Meer Y, Veerman RM. What Autocorrelation Tells Us about Motor Variability: Insights from Dart Throwing. PLoS ONE. 2013;8(5):e64332. doi: https://doi.org/10.1371/journal.pone.0064332.

75. van Beers RJ, Brenner E, Smeets JBJ. Random walk of motor planning in task-irrelevant dimensions. J Neurophysiol. 2013;109(4):969-77. doi: https://doi.org/10.1152/jn.00706.2012.

76. van Beers RJ. How Does Our Motor System Determine Its Learning Rate? PLoS ONE. 2012;7(11):e49373. doi: https://doi.org/10.1371/journal.pone.0049373.

77. van der Kooij K, Brenner E, van Beers RJ, Smeets JBJ. Visuomotor Adaptation: How Forgetting Keeps Us Conservative. PLoS ONE. 2015;10(2):e0117901. doi: 10.1371/journal.pone.0117901.

78. López-Moliner J, Vullings C, Madelain L, van Beers RJ. Prediction and final temporal errors are used for trial-to-trial motor corrections. Scientific Reports. 2019;9(1):19230. doi: 10.1038/s41598-019-55560-6.

79. Coltman SK, Beers RJv, Medendorp WP, Gribble PL. Sensitivity to error during visuomotor adaptation is similarly modulated by abrupt, gradual, and random perturbation schedules. J Neurophysiol. 2021;126(3):934-45. doi: 10.1152/jn.00706.2021. PubMed PMID: 34379553.

80. Tumer EC, Brainard MS. Performance variability enables adaptive plasticity of ‘crystallized’ adult birdsong. Nature. 2007;450(7173):1240-4. doi: https://doi.org/10.1038/nature06390.

81. Sober SJ, Brainard MS. Adult birdsong is actively maintained by error correction. Nat Neurosci. 2009;12(7):927-31. doi: https://doi.org/10.1038/nn.2336.

82. Veit L, Tian LY, Monroy Hernandez CJ, Brainard MS. Songbirds can learn flexible contextual control over syllable sequencing. eLife. 2021;10:e61610. doi: https://doi.org/10.7554/eLife.61610.

83. Hauser MD, Chomsky N, Fitch WT. The faculty of language: what is it, who has it, and how did it evolve? Science. 2002;298(5598):1569-79. Epub 2002/11/26. doi: 10.1126/science.298.5598.1569. PubMed PMID: 12446899.

84. Grafton ST. Malleable templates: reshaping our crystallized skills to create new outcomes. Nat Neurosci. 2008;11(3):248-9. doi: https://doi.org/10.1038/nn0308-248.

85. Daley MA, Biewener AA. Running over rough terrain reveals limb control for intrinsic stability. Proc Natl Acad Sci USA. 2006;103(42):15681-6. doi: https://doi.org/10.1073/pnas.0601473103.

86. Othayoth R, Thoms G, Li C. An energy landscape approach to locomotor transitions in complex 3D terrain. Proc Natl Acad Sci USA. 2020;117(26):14987-95. doi: 10.1073/pnas.1918297117.

87. Jindrich DL, Full RJ. Many-Legged Maneuverability: Dynamics of Turning in Hexapods. J Exp Biol. 1999;202(12):1603-23. doi: https://doi.org/10.1242/jeb.202.12.1603.

88. Full RJ, Kubow T, Schmitt J, Holmes P, Koditschek DE. Quantifying dynamic stability and maneuverability in legged locomotion. Integrative and Comparative Biology. 2002;42(1):149-57. doi: https://doi.org/10.1093/icb/42.1.149.

89. Gallivan JP, Chapman CS, Wolpert DM, Flanagan JR. Decision-making in sensorimotor control. Nat Rev Neurosci. 2018;19(9):519-34. Epub 2018/08/10. doi: 10.1038/s41583-018-0045-9. PubMed PMID: 30089888; PubMed Central PMCID: PMCPMC6107066.

90. Nashed JY, Crevecoeur F, Scott SH. Rapid Online Selection between Multiple Motor Plans. The Journal of Neuroscience. 2014;34(5):1769-80. doi: 10.1523/jneurosci.3063-13.2014.

91. Wong AL, Haith AM, Krakauer JW. Motor Planning. The Neuroscientist. 2015;21(4):385-98. doi: 10.1177/1073858414541484. PubMed PMID: 24981338.

92. Zeni JA, Richards JG, Higginson JS. Two simple methods for determining gait events during treadmill and overground walking using kinematic data. Gait Posture. 2008;27(4):710-4. doi: http://dx.doi.org/10.1016/j.gaitpost.2007.07.007.

93. Donelan JM, Kram R, Kuo AD. Mechanical and metabolic determinants of the preferred step width in human
walking. Proc Roy Soc B Biol Sci. 2001;268(1480):1985-92. doi: https://doi.org/10.1098/rspb.2001.1761.

94. Wang Y, Srinivasan M. Stepping in the direction of the fall: the next foot placement can be predicted from current upper body state in steady-state walking. Biol Lett. 2014;10(9):20140405. doi: http://dx.doi.org/10.1098/rsbl.2014.0405.

95. Donelan JM, Shipman DW, Kram R, Kuo AD. Mechanical and metabolic requirements for active lateral stabilization in human walking. J Biomech. 2004;37(6):827-35. doi: https://doi.org/10.1016/j.jbiomech.2003.06.002.

96. McAndrew PM, Wilken JM, Dingwell JB. Dynamic Stability of Human Walking in Visually and Mechanically Destabilizing Environments. J Biomech. 2011;44(4):644-9. doi: http://dx.doi.org/10.1016/j.jbiomech.2010.11.007.

97. McAndrew Young PM, Wilken JM, Dingwell JB. Dynamic Margins of Stability During Human Walking in Destabilizing Environments. J Biomech. 2012;45(6):1053-9. doi: http://dx.doi.org/10.1016/j.jbiomech.2011.12.027.

98. Rankin BL, Buffo SK, Dean JC. A Neuromechanical Strategy for Mediolateral Foot Placement in Walking Humans. J Neurophysiol. 2014;112(2):374-83. Epub April 30, 2014. doi: https://doi.org/10.1152/jn.00138.2014.

99. Stimpson KH, Heitkamp LN, Horne JS, Dean JC. Effects of walking speed on the step-by-step control of step width. J Biomech. 2018;68:78-83. doi: https://doi.org/10.1016/j.jbiomech.2017.12.026.

100. Ghahramani Z, Wolpert DM. Modular Decomposition in Visuomotor Learning. Nature. 1997;386(6623):392-5. doi: https://doi.org/10.1038/386392a0.

101. Efron B, Tibshirani RJ. An Introduction to the Bootstrap (1st Ed.). Chapman and Hall/CRC. 1994. doi: https://doi.org/10.1201/9780429246593.