The Glycemic Index of Sport Nutrition Bars Affects Performance and Metabolism During Cycling and Next-Day Recovery

by
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Low-glycemic index carbohydrates are potentially better for endurance performance as they result in greater fat oxidation and lower carbohydrate oxidation due to lower insulin release. We compared the effects of pre-exercise feeding with a low-glycemic index lentil-based sports nutrition bar, a commercially-available sports nutrition bar with moderate-glycemic index, and a non-caloric placebo on metabolism and performance during endurance cycling (Trial 1). Using a randomized, counterbalanced, crossover design, endurance-trained individuals (n = 11; eight males; 26 ± 6y; VO2peak 51.4 ± 1.6 mL/kg/min) consumed 1.5 g/kg available carbohydrate from a lentil bar and a moderate-glycemic index bar, as well as a placebo, 1h before endurance cycling (75 min at 65% VO2peak, followed by a 7 km time trial). We also compared post-exercise consumption of the low-glycemic index bar with another moderate-glycemic index bar on next-day exercise performance as an assessment of recovery (Trial 2). In Trial 1, fat or carbohydrate oxidation rates were not different between the bar conditions (p > 0.05). Blood lactate was lower during the low- versus the moderate-glycemic index condition after 75 minutes of cycling (2.6 versus 4.0 mmol/L, p < 0.05) and at the end of the time trial (7.4 versus 9.1 mmol/L, p < 0.05). Time trial performance improved (p < 0.05) after consumption of the low- (574 ± 55 s) and moderate-glycemic index (583 ± 59 s) bars compared to the placebo (619 ± 81 s). In Trial 2 (next-day recovery), performance improved (p < 0.05) with the low-glycemic index bar (547 ± 42 s) compared to the moderate-glycemic index bar (569 ± 42 s) and the placebo (566 ± 34 s). Low- and moderate-glycemic index sports nutrition bars improved cycling exercise performance; however, only the low-glycemic index bar improved next day performance.

Key words: carbohydrate, exercise, fatigue, endurance.

Introduction

Carbohydrate consumption is important for endurance performance and recovery from endurance exercise (Litte et al., 2009a). Carbohydrates with a low glycemic index (GI) result in slow release of glucose into the bloodstream after digestion (Little et al., 2009a). This may be of benefit if consumed before endurance exercise since low-GI carbohydrates would supply a sustained source of energy to contracting muscle (Little et al., 2009b). The low insulin response with consumption of low-GI carbohydrates may also be beneficial for substrate metabolism because high insulin levels inhibit fat lipolysis and oxidation (Horowitz et al., 1997) and, therefore, promote carbohydrate oxidation. Thus, the consumption of low-GI carbohydrates may also promote sparing of muscle glycogen during endurance exercise and improve performance (Little et al., 2010).

Whether lower-GI carbohydrates can actually provide greater benefit for exercise performance compared to higher-GI carbohydrates is an area of controversy. Two recent meta-analyses had contrasting conclusions, with one showing a statistically significant benefit...
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(p=0.001) of pre-exercise lower-GI meals/beverages for improving performance over high GI meals/beverages (Heung-Sang Wong et al., 2017) and the second showing no differences in performance between GI conditions (Burdon et al., 2017). Each meta-analysis used similar criteria for their included studies; however, these studies differed slightly between each meta-analysis. The meta-analysis methods differed with one using a more conservative model (i.e. random effects; Burdon et al., 2017) and one using a more liberal model (i.e. fixed effects), which is more likely to result in a statistically significant difference (Heung-Sang Wong et al., 2017). Most importantly the review by Burdon et al. (2017) performed a separate meta-analysis for each type of performance outcome (i.e. time trial, sub-maximal exercise followed by a time trial, and time-to-exhaustion); whereas the review by Heung-Sang Wong et al. (2017) standardized all performance outcomes and combined them into one large meta-analysis. The latter approach includes more studies per meta-analysis and therefore has greater statistical power, which could account for their finding of statistically significant performance benefits for lower-GI versus higher-GI carbohydrates.

Lentils have a very low GI and when consumed before endurance exercise they enhance metabolism (i.e. fat oxidation is increased, carbohydrate oxidation is reduced) compared to other carbohydrate sources (Bennett et al., 2012; Little et al., 2009b). This is most likely due to the smaller release of insulin after a low-GI meal because lower insulin levels result in greater fatty acid oxidation (Horowitz et al., 1997). Consuming boiled lentils is not very palatable (Little et al., 2009a, b); therefore, we sought to determine whether lentils incorporated into a sport nutrition bar would offer similar benefits. Our objective was to determine whether a low-GI lentil-based sport nutrition bar consumed prior to exercise could optimise metabolism and exercise performance compared to a sport nutrition bar with a moderate GI.

Moderate to high-GI carbohydrates are generally promoted for recovery from endurance exercise because they enhance glycogen re-synthesis after endurance exercise, most likely because they stimulate greater insulin release, which promotes glucose uptake into muscle (Burke et al., 1993). However, when meals of different GIs are compared during recovery after strenuous exercise, studies indicate either superiority of low-GI meals or no difference between meals of different GI on subsequent exercise performance (Brown et al., 2013; Erith et al., 2006; Stevenson et al., 2005). Low-GI meals may promote either greater sparing of glycogen during the recovery exercise session or greater sparing of other important intra-muscular substrates, such as intra-muscular triglycerides allowing for superior performance during the recovery exercise session (Stevenson et al., 2009; Trenell et al., 2008). We hypothesized a low-GI lentil-based sports nutrition bar consumed before exercise would enhance fat oxidation, reduce carbohydrate oxidation, and enhance endurance performance compared to a moderate-GI bar and that a low GI bar consumed during recovery from exhaustive exercise would be superior to a moderate-GI bar for improving next-day exercise performance.

Methods

Participants

Eight males and three females with a moderate level of cycling experience were recruited (26 ± 6 y, 178 ± 3 cm, 72.9 ± 4.0 kg, and VO2peak 51.4 ± 1.6 mL/kg/min). The study followed ethical guidelines consistent with the Declaration of Helsinki; The University of Saskatchewan Research Ethics Board approved the study protocol, and participants provided signed informed consent.

Measures, Design, and Procedures

On their first visit to the laboratory, participants completed a cycle ergometer (Monark, Ergomedic 874 E, Sweden) protocol to determine peak oxygen consumption (VO2peak). Participants were instructed to pedal at 80 rpm. After a 5-min warm-up, the initial resistance was set at 0.5 kp for females and 1-1.5 kp for males, then increased every 2 minutes by 0.5 kp until the participant could not keep a pedaling cadence of 70 rpm. Throughout testing, VO2 was recorded by open circuit indirect calorimetry (Vmax Series 29 Calorimeter, SensorMedics, USA). The heart rate was measured continuously using a Polar 610i heart rate monitor (Polar Electro Oy, Kempele, Finland). Criteria that were used for determining VO2peak included achieving a plateau in VO2 respiratory exchange rate more than 1.15, and 95
percent of the age-predicted maximum heart rate (Howley et al., 1995). Following the VO2peak test, participants were familiarized with performance testing to be used in the study. This involved a 7 km time trial where participants cycled as fast as possible at an intensity corresponding to 65% VO2peak. During this familiarization test and all performance tests subsequently used in the study, participants were blinded to the elapsed time, speed, and cadence by masking the cycle display screen. Participants were allowed to view their elapsed distance during the test. A week later, participants performed a familiarization test on the cycle ergometer that simulated the complete testing protocol. This involved cycling at an intensity corresponding to 65% VO2peak for 75 minutes, followed by a 7 km time trial, similar to the first visit. The exercise test was designed to simulate the distance that cyclists would typically cover during an Olympic-distance triathlon (i.e. 40 km).

Using a randomized, counterbalanced, cross-over design, participants consumed 1.5 g/kg available carbohydrate from a low-GI lentil bar (the Genki Bar; Genki Bar Foods Inc., Winnipegosis, Manitoba, Canada), a moderate-GI bar (PowerBar Harvest Inc., Berkeley, California) or consumed a placebo (Diet Flavored Jell-O) after an overnight fast, and one hour before endurance cycling for 75 minutes at an intensity that corresponded to 65% VO2peak for 75 minutes, followed by a 7 km time trial, similar to the first visit. The exercise test was designed to simulate the distance that cyclists would typically cover during an Olympic-distance triathlon (i.e. 40 km).

We previously evaluated the GI of all three bars and determined the lentil bar to have a low GI of 47, which was significantly lower than the Harvest PowerBar™ (GI = 56) and the PowerBar™ Recovery (GI = 58) (Chilibeck et al., 2011). Both Powerbars™ were classified as having a "moderate" GI. Each feeding condition was separated by at least one week. Females performed each condition one month apart so that they were always in the same phase of their menstrual cycle. The individual administering the exercise tests was blinded to all feeding conditions by having a separate researcher, who was not involved in data collection to prepare the bars or the placebo before the tests and having the participant consume the bars or placebo in a separate room than the researcher performing the exercise tests. The participants were blinded to the bar conditions by removal of the bar wrappers; however, most participants could discern the difference between the bars based on their appearance, especially the PowerBar™ Recovery because it is coated in chocolate, unlike the other bars. Participants were unaware of the hypotheses of the study.

Respiratory gases were collected from the 10th to the 20th minute and the 60th to the 70th minute during the 75 minutes of constant-load cycling to estimate fat and carbohydrate oxidation. Oxygen consumption (VO2), expired carbon dioxide (VCO2) and Respiratory Exchange Ratio (RER) were calculated breath-by-breath using open-circuit indirect calorimetry (Vmax Series 29 Calorimeter; Sensor Medics). Carbohydrate and fat oxidation were based on calculations for moderate to high-intensity exercise (Jeukendrup and Wallis, 2005). Rating of perceived exertion (RPE) was measured by using the modified Borg scale (i.e. scale of 1 to 10) at the 25th, 50th, and 75th minute of cycling and at the end of the time trial (Borg, 1973).

Capillary blood was drawn from the fingertip and assessed using two different meters (AccuCheck Compact Plus and Accutrend GC; Roche Diagnostics, Sarstedt, Nümbrecht, Germany) to measure glucose and lactate levels, respectively, at the 5th, 15th, 30th, and 60th min after consuming nutrition bars or the placebo, and after 25, 50, and 70 min cycling and at the end of the 7 km time trial. Venous blood samples from an antecubital vein were collected at three times (i.e., an hour after consumption of the nutrition bars or placebo, after 75 min cycling, and following the 7 km time trial) for assessment of non-esterified fatty acids.
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(NEFA), insulin, and catecholamines. The blood samples were maintained in 10 mL tubes (BD Vacutainer SST, Company, City) for 30 minutes to clot. Serum was separated by centrifugation for 10-12 minutes at 3500 rpm and stored at -80°C. Insulin, epinephrine, and norepinephrine were determined using an ELISA according to the manufacturer’s directions (Insulin El, Alpco Diagnostics, City; 2-cat EIA, Alpco Diagnostics). The serum NEFA assay was performed using a protocol with an oleic acid standard solution as per the manufacturer’s directions (NEFAHR (2), Wako Diagnostics). All assays for an individual participant were performed on the same plate to eliminate inter-assay variability. The intra-assay CVs for insulin, catecholamines, and NEFA were < 10%.

To minimize any possible influences of diet, physical activity, and sleep variability on study measurements, participants were asked to complete a 24-hour diet, physical activity, and sleep record the day before the first condition and the day of the first condition. The logs were photocopied and returned to participants. Each participant was instructed to repeat the same diet, physical activity, and sleep pattern for the subsequent conditions. Water intake during the exercise tests was matched across conditions. The participant was allowed to drink as much water as they wanted during the first condition and then given the same amount of water during the other conditions.

Statistical Analysis

All statistical analyses were performed using SPSS for McIntosh (SPSS Version 20; SPSS Inc., Delaware). A two-factor repeated-measures ANOVA (meal and time) was used to assess differences in blood glucose, blood lactate, RER, carbohydrate oxidation, fat oxidation, serum insulin, NEFA, catecholamines (i.e. epinephrine and nor-epinephrine) and RPE. A one factor (meal) repeated measures ANOVA was used to assess time trial performance for the first and second day (i.e. recovery day) of testing. Where significance was noted, an LSD post hoc test was used to determine differences between pairs of means. The level of significance was set at $p < 0.05$. Results are presented as means ± SD.

Results

Both sports bar conditions elicited a significantly better time trial performance on day 1 than the placebo condition ($p = 0.001$; Figure 1A). The sport bar conditions were not different from each other. The next-day recovery time trial performance was significantly improved in the low-GI condition compared to the moderate-GI bar and the placebo ($p = 0.006$; Figure 1B).

A trend for a condition main effect for RPE was present ($p = 0.069$; low-GI bar = 6.3 ± 0.7, moderate-GI bar = 6.4 ± 0.8; placebo = 6.7 ± 0.6). There was no meal x time interaction. As expected, a significant time main effect ($p < 0.001$) was observed with RPE increasing during the exercise (data not shown).

Plasma glucose between low-GI and moderate-GI conditions did not differ throughout the postprandial period and exercise time. As expected, plasma glucose was significantly lower in the placebo condition compared to the low-GI and moderate-GI conditions at all time-points during the postprandial period ($p = 0.0001$) (Figure 2A).

Carbohydrate and fat oxidation rates were similar in the low- and moderate-GI conditions. Carbohydrate oxidation was lower ($p = 0.02$) and fat oxidation higher ($p = 0.0001$) in the placebo condition versus both sport nutrition bars (Table 2).

No significant difference in the respiratory exchange ratio was apparent between the nutrition bar conditions; however, the respiratory exchange ratio was lower in the placebo condition versus sports bar conditions ($p = 0.0001$; Table 2).

There was a condition by time interaction, where the insulin level during the low-GI condition after 75 min cycling was significantly lower than the insulin level in the moderate-GI condition ($p = 0.025$). The insulin response with the placebo was lower than the other conditions at the start of exercise and after the 7 km time trial, and was lower than the moderate-GI condition after 75 min cycling ($p = 0.005$) (Table 3).

NEFA levels were significantly higher after 75 min cycling and the 7 km time trial compared to the onset of cycling across the three conditions (time main effect; $p = 0.0001$). Also NEFA concentrations were significantly higher in the placebo condition versus the low-GI and moderate-GI conditions across time points (condition main effect; $p = 0.002$, Table 3).

The concentrations of epinephrine and norepinephrine were similar in all experimental
conditions across time points. There was a time main effect, with catecholamines increasing over the exercise test ($p = 0.0001$, Table 3).

| Table 1 | Characteristics of the nutrition bars for a 70 kg participant |
| --- | --- | --- | --- |
| Description | Low GI bar | Moderate GI (Recovery) | Moderate GI bar |
| Energy (kcal) | 830 | 756 | 910 |
| Fat (g) | 19 | 17 | 35 |
| CHO total (g) | 127 | 119 | 105 |
| CHO available (g) | 105 | 105 | 105 |
| Protein (g) | 39 | 35 | 42 |
| Glycemic index | 47 | 56 | 58 |

| Table 2 | Substrate oxidation and respiratory exchange ratio (RER) during cycling |
| --- | --- | --- | --- |
| Time | Low GI bar | Moderate GI bar | Placebo* |
| 10 – 20 min | 60 – 70 min | 10 – 20 min | 60 – 70 min | 10 – 20 min | 60 – 70 min |
| Fat Oxidation (g/min) | 0.23 ± 0.2 | 0.46 ± 0.6 | 0.27 ± 0.1 | 0.47 ± 0.2 | 0.42 ± 0.2 | 0.7 ± 0.2 |
| Carbohydrate oxidation (g/min) | 2.25 ± 0.6 | 1.83 ± 0.5 | 2.15 ± 0.7 | 1.87 ± 0.6 | 1.85 ± 0.5 | 1.31 ± 0.6 |
| RER | 0.94 ± 0.04 | 0.89 ± 0.03 | 0.93 ± 0.04 | 0.89 ± 0.03 | 0.89 ± 0.03 | 0.83 ± 0.01 |

* Significantly different in the placebo condition versus the nutrition bars conditions for all variables across all time points ($p<0.05$). Values are mean ± SD.

| Table 3 | Insulin, non-esterified fatty acids (NEFA), epinephrine, and norepinephrine concentrations in the three conditions. |
| --- | --- | --- | --- | --- |
| Condition | Onset of cycling | After 7 km | After 7 km | After 7 km |
| Low GI bar | Moderate GI bar | Placebo |
| Insulin (mmol/L)$^a$ | 30 ± 16$^a$ | 8 ± 9$^b$ | 17 ± 27$^a$ | 36 ± 24$^a$ | 15 ± 17 | 16 ± 17$^a$ | 6 ± 5 | 7 ± 11$^b$ | 3 ± 2 |
| NEFA$^{cd}(mmol/L)$ | 0.29 ± 0.07 | 0.67 ± 0.08 | 0.48 ± 0.07 | 0.24 ± 0.07 | 0.7 ± 0.07 | 0.57 ± 0.07 | 0.4 ± 0.07 | 0.4 ± 0.07 | 1 ± 0.4 | 0.87 ± 0.3 |
| Epinephrine (nmol/L)$^e$ | 0.1 ± 0.03 | 0.2 | 0.01 | 0.4 | 0.2 ± 0.03 | 1.3 ± 0.03 | 0.2 ± 0.03 | 0.8 ± 0.03 | 1.2 ± 0.03 | 1.3 ± 0.3 |
| Norepinephrine (nmol/L)$^e$ | 7.3 ± 0.7 | 15.5 ± 1.4 | 28 ± 2.0 | 8.1 ± 4.9 | 13.7 ± 2.0 | 26.4 ± 2.0 | 8 ± 4 | 16.2 ± 6.8 | 21.1 ± 10.2 |

Values are mean ± SD of participants.

$^a$Different compared to the placebo ($p < 0.01$)

$^b$Different compared to the moderate-GI condition

$^c$Time main effect, 75 min and post-time 7 km time trial different than onset of cycling ($p < 0.01$)

$^d$Condition main effect, placebo different than low- and moderate-GI bar across time points ($p < 0.01$)

$^e$Time main effect, values for epinephrine and norepinephrine increased across all time points ($p < 0.01$)
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Figure 1
A) Day 1 time trial performance in the three conditions. B) Time trial performance for the day after exercise (recovery day) in the three conditions. Values are mean and error bars are SD.
Figure 2

A) Plasma glucose concentration and B) lactate vs. time, at: baseline, postprandial and during exercise.

Values are mean and error bars are SD.

Blood glucose was lower in the placebo compared to the bar conditions (p<0.01)

*Blood lactate was higher in the moderate-GI condition versus the low-GI condition (p<0.01)

**Blood lactate was higher in the moderate-GI condition versus the low-GI and placebo conditions (p<0.01)
The data shown in Figure 2B illustrate the impact of the low-GI and moderate-GI bars and the placebo on blood lactate levels during the cycling test and after the 7 km time trial. A condition by time interaction was present with blood lactate levels significantly higher after 75 min of exercise in the moderate-GI compared to the low-GI condition \((p = 0.003)\). Blood lactate was also higher after the 7 km time trial in the moderate-GI compared to low-GI and placebo conditions \((p = 0.003)\).

All participants complied with instructions to follow similar dietary, sleep, and physical activity patterns across conditions. This was done by checking the initial log completed on the first condition with participants on subsequent conditions. Timing of meals and meal composition before exercise tests and during recovery were similar across conditions with mean daily energy intake of 2465 (SD 860) kcal, protein intake of 109 (SD 46) g, carbohydrate intake of 305 (SD 117) g, and fat intake of 88 (SD 39) g.

**Discussion**

The most important finding of this study was that the consumption of bars of differing GI improved performance compared to the placebo when consumed an hour before exercise; however, there was no difference between the bars of a different GI. Consumption of a low-GI bar during recovery was most effective for improving next-day performance. To the best of our knowledge, this study is the first to assess the effect of sport nutrition bars of different GIs on exercise performance, metabolic responses to exercise, and recovery from exercise. This is important given that many sports nutrition bars are commercially available promoting improved exercise performance and recovery.

The consumption of low-GI compared to moderate- or high-GI carbohydrates results in lower and more stable glucose and insulin response. Therefore, low-GI foods could be superior for stimulating fat oxidation and reducing carbohydrate oxidation during exercise because insulin inhibits fat oxidation (Horowitz et al., 1997). The blood glucose response to the moderate-GI bar in our study, however, was not significantly different than the low-GI bar (Figure 2a), which differed from our previous determination of a significantly higher glucose response in the moderate-GI bar (Chilibeck et al., 2011). The differences in findings could be associated with the more precise assessment of the GI in our previous study which evaluated multiple glucose measurements across each time point and was based on the glucose area under the curve over 2 hours. The current study only assessed glucose once at each time point and assessed glucose response for one hour. Although GIs of the nutrition bars were significantly different, the GIs (47 versus 56) may not have been different enough to elicit significantly different metabolic responses. The insulin response to the two bars was similar, with the moderate-GI bar eliciting a higher insulin response only at the 75th minute mark of exercise.

A lack of difference in fat and carbohydrate oxidation, mobilization of fatty acids, and catecholamine response between the bar conditions was different from other studies comparing low-GI lentils to high-GI meals (Bennett et al., 2012; Little et al., 2009b; Moore et al., 2010). Again this may be due to the relatively small difference in GIs between the bars. The differences in the GI between the bars in the current study was only 9 GI units; whereas the difference in the GI of other studies detecting differences in substrate oxidation had differences ranging from 36 to 91 GI units for lower versus higher-GI conditions (Bennett et al., 2012; Little et al., 2009b). Males and females respond differently regarding fat oxidation during endurance exercise (Tarnopolsky, 2008) and therefore the inclusion of females may have distorted our data. A limitation of our study is the inclusion of only a small number of females, which did not allow us to include sex as a factor in our analysis; however, when we excluded females from our analysis, there were still no differences between GI conditions for substrate oxidation.

Lactate production late in the exercise session was lower during the low-GI condition (Figure 2b), which may have indicated a smaller use of glycogen during anaerobic metabolism. This is consistent with other studies showing a higher blood lactate concentration during exercise when high- compared to low-GI foods are consumed before exercise (Karamanolis et al., 2011; Wee et al., 1999, 2005). Wee et al. (2005) suggested that a greater accumulation of muscle and blood lactate during treadmill running was accompanied by reductions of muscle glycogen during a high-GI
trial. In contrast with this, other studies failed to find any significant difference in lactate accumulation when food with different GIs was consumed prior to continuous exercise (Kirwan et al., 1998; Thomas et al., 1991).

A lower use of glycogen through anaerobic metabolism may have contributed to the superior performance during the low-GI bar condition versus the moderate-GI condition during the next-day performance. In agreement with this, Stevenson et al. (2009) reported that a low-GI diet consumed in the 24 h period following continuous heavy exercise led to a greater endurance capacity during steady state exercise in the post-absorptive state the following day compared with a high-GI carbohydrate diet. Moreover, Trenell et al. (2008) showed that using a low-GI recovery diet may spare intramuscular triglyceride utilised during recovery exercise performance, whereas a high-GI diet reduced NEFA availability throughout exercise and increased reliance on intramuscular lipids as an energy source during moderate intensity exercise. Intramuscular triglyceride, in addition to glycogen, may be an important energy source for skeletal muscle during endurance exercise.

One of the unique aspects of this study was testing of different sport nutrition bars for the subsequent exercise metabolism and performance. Consuming nutrition bars has become popular among athletes since they are convenient and well portion-controlled (Rauch et al., 1999). In the present study we provided multiple nutrition bars instead of a single bar in order to meet recommended carbohydrate intake prior to endurance exercise. To the best of our knowledge, this study was the first to investigate the effect of consuming multiple nutrition bars on exercise performance; whereas, most previous nutritional bar studies failed to show beneficial results because the amount of bars consumed was too few based on previous research of how much carbohydrate is needed to improve performance (Kolkhorst et al., 1998; Oliver and Tremblay, 2002; Rauch et al., 1999). A limitation of the study was that changes in the muscular glycogen and intramuscular triglyceride stores were not measured following the first time trial and the subsequent time trial the next day to disclose the possible metabolic reasons for better performance in the low-GI condition.

In summary, low- and moderate-GI bars elicited similar metabolic responses and had similar benefits for endurance performance; however, a low-GI bar enhanced recovery as indicated by superior next-day exercise performance. This might be attributable to lower anaerobic glycogenolysis and, therefore, sparing of muscle glycogen late in the initial exercise session. Irrespective of the mostly similar responses in the current study to low- and moderate-GI bars, consumption of low-GI bars by athletes may have long-term nutritional health benefits compared to those bars that elicit an undesirable glucose response (Tegelman et al., 1996).

**Practical Recommendations**

Low or moderate-GI sport nutrition bars, when consumed in adequate amounts to meet guidelines for pre-exercise carbohydrate consumption (i.e. ≥ 1.0 g/kg available carbohydrate) are beneficial for improving endurance exercise performance. When consumed during recovery, low-GI sport nutrition bars appear to provide the greatest benefit with regard to the next-day endurance exercise performance.

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