Carbohydrate Hydrogel Products Do Not Improve Performance or Gastrointestinal Distress During Moderate-Intensity Endurance Exercise

Andy J. King  Joshua T. Rowe
Australian Catholic University  University of Leeds

Louise M. Burke
Australian Catholic University and Australian Institute of Sport

The benefits of ingesting exogenous carbohydrate (CHO) during prolonged exercise performance are well established. A recent food technology innovation has seen sodium alginate and pectin included in solutions of multiple transportable CHO, to encapsulate them at pH levels found in the stomach. Marketing claims include enhanced gastric emptying and delivery of CHO to the muscle with less gastrointestinal distress, leading to better sports performance. Emerging literature around such claims was identified by searching electronic databases; inclusion criteria were randomized controlled trials investigating metabolic and/or exercise performance parameters during endurance exercise >1 hr, with CHO hydrogels versus traditional CHO fluids and/or noncaloric hydrogels. Limitations associated with the heterogeneity of exercise protocols and control comparisons are noted. To date, improvements in exercise performance/capacity have not been clearly demonstrated with ingestion of CHO hydrogels above traditional CHO fluids. Studies utilizing isotopic tracers demonstrate similar rates of exogenous CHO oxidation, and subjective ratings of gastrointestinal distress do not appear to be different. Overall, data do not support any metabolic or performance advantages to exogenous CHO delivery in hydrogel form over traditional CHO preparations; although, one study demonstrates a possible glycogen sparing effect. The authors note that the current literature has largely failed to investigate the conditions under which maximal CHO availability is needed; high-performance athletes undertaking prolonged events at high relative and absolute exercise intensities. Although investigations are needed to better target the testimonials provided about CHO hydrogels, current evidence suggests that they are similar in outcome and a benefit to traditional CHO sources.

Keywords: encapsulated carbohydrate, glycogen, gut, sports nutrition, oxidation, sports drink

Recent interest in the 2-hr marathon (Caesar, 2019) has focused attention on an important sports nutrition strategy; consumption of carbohydrate (CHO) during exercise to contribute to the substantial fuel costs of some endurance events. Events of sufficient intensity and duration to be limited by CHO availability benefit from an exogenous CHO supply (Stellingwerff & Cox, 2014), with mechanisms including fuel provision once muscle glycogen is depleted (Coyle et al., 1986), spared liver (Gonzalez et al., 2015; Wallis et al., 2006) and muscle (King et al., 2018; Tsintzas et al., 1995, 1996) glycogen use, and central nervous system benefits (Burke & Maughan, 2015). A sliding scale of intake is recommended, according to event fuel needs and specific mechanisms underpinning performance benefits (Thomas et al., 2016). Upper targets for fuel-demanding events (80–90+ g·hr\(^{-1}\) CHO), which aim to maximize the contribution of exogenous CHO to substrate use, are often challenged by the ability to consume, tolerate, and absorb large amounts of CHO (de Oliveira & Burini, 2014). Factors include the availability of foods/drinks to meet CHO targets in practical amounts/volumes, the effect of the mode and intensity of exercise on gastrointestinal (GI) comfort and function (de Oliveira & Burini, 2009), the role of specific “gut training” (Cox et al., 2010), and characteristics of the CHO source. Here it has been shown that the use of CHO blends (“multiple transportable CHO” such as glucose [G] and fructose [F]) can maximize gut uptake via the use of different intestinal transport mechanisms, assisting with substrate delivery and the management of gut comfort (Jeukendrup, 2010).

Recently, specialized sports foods claiming to address such factors via the use of “hydrogel technology” have become commercially available (Sutehall et al., 2018). These supplements, combining typical CHO sources with pectin (a soluble fiber)
and alginate (a polymer derived from seaweed) undergo gelation on contact with low pH solutions, such as stomach acid, to encapsulate the CHO (Marciani et al., 2019). Enhanced rates of gastric emptying could deliver this “hydrogel” to the small intestine where it dissolves in the higher pH environment for absorption, leading to reduced gut discomfort, enhanced muscle CHO delivery, and performance benefits (Figure 1). Indeed, according to testimonials, the commercial product has been quickly adopted by elite athletes (Sutehall et al., 2018) and publicized in sporting successes including the 1:59 marathon project, leading to marketing claims that it is the “world’s fastest sports fuel” (Maurten, 2020). Noting that this has largely occurred in the absence of scientific validation of these claims, we undertook a review of newly published investigations of hydrogel CHO to determine whether they achieve better GI characteristics, substrate delivery, and performance effects under exercise conditions than traditional sports drinks and gels.

### Methods

A search of electronic databases for studies published up to May 14, 2020 was independently completed by two authors (A. King/J. Rowe), with the key methodological process and considerations involved in including/excluding data summarized in Figure 2.

To be eligible for this review, studies were required to have investigated a CHO hydrogel compound during prolonged, endurance exercise defined as continuous running, cycling, triathlon, rowing, swimming, and cross-country skiing greater than 1-hr duration. Studies with exercise durations lasting 1 hr or less were excluded because CHO ingestion is unlikely to be beneficial during shorter duration exercise (Burke et al., 2011; Thomas et al., 2016). CHO mouth rinse studies were also excluded as the primary mechanism whereby performance is improved is neurological in origin.

The CHO hydrogel used could be a commercially available product or a laboratory-manufactured solution, provided that the active substance included as a gelling agent was known to encapsulate ingested CHO in the stomach. A control comparison/condition was required to be a typical CHO control, matched for CHO dose and type, or a placebo. Studies were included if they reported data on one or more of physiological or performance variables (Table 1). Review articles and case studies were excluded.

From the available studies, between-condition differences for hydrogel and comparison products were calculated. Standardized effect sizes (ES) were calculated using Hedge’s g adjustment for small samples with 95% confidence intervals for the ES computed. No statistical adjustments for ES were made for carryover effects, since suitable washout periods were included in these crossover trials. No assessment of publication bias was undertaken with the low number of studies and the likely presence of heterogeneity; we noted that this can affect the robustness of publication bias analysis (Ioannidis & Trikalinos, 2007). Forest plots were produced to provide a visual comparison of effects in studies measuring exercise performance.

### Results

We located six studies (Table 1) which compared a CHO hydrogel containing alginate and/or other gelling compounds with a noncaloric hydrogel placebo (n = 1) or CHO fluids of matched CHO (n = 5). Although CHO hydrogels were similar in composition, containing a mixture of maltodextrin (MD) and F, total CHO ranged from 68 to 132 g·hr⁻¹. Five of the six studies included a matched condition for dose and type of CHO (i.e., MD + F fluid) and two studies included a G or MD only fluid matched for CHO dose. Therefore, four studies met guidelines for upper range CHO.

---

**Figure 1** — Mechanisms of CHO hydrogel formation and delivery to the small intestine. Despite benefits to gastric emptying with hydrogel-encapsulated CHO, the rate-limiting step of exogenous CHO oxidation lies in the intestinal transport of monosaccharides. CHO = carbohydrate.
“CHO-rich” meal (Flood et al., 2020). The only study undertaken under fasting conditions (Barber et al., 2020) focused on exogenous CHO oxidation as the primary outcome, which is not affected by prior muscle glycogen content (Margolis et al., 2019). Between-conditions comparisons remain valid despite the difference in breakfast protocol to other studies.

**Exercise Performance**

Performance was similar with MD + F hydrogel, isocaloric MD + F fluids (Figure 3; Baur et al., 2019; Flood et al., 2020; McCubbin et al., 2019; Mears et al., 2020b), or noncaloric hydrogel (Pettersson et al., 2019). Relative performance changes between CHO hydrogel and fluids ranged between +1.05% and +3.8% but were not statistically significant (p <.05). The largest change reported by Mears et al. (2020b) during a ~20-min fixed-work time trial was a moderately higher workload (3.8% improvement, ES = 0.51) with CHO hydrogel ingestion compared with the dose matched MD + F control. All other performance effects were very small (ES < 0.10).

**Physiological Measures**

Exogenous CHO oxidation was measured in two studies, but only one included a comparative CHO condition (Barber et al., 2020) with exogenous CHO oxidation peaking at 1.1 ± 0.3 g·min⁻¹ in both the MD + F hydrogel and MD + F fluid (Table 1). MD + G ingestion resulted in a moderately lower oxidation rate (0.92 ± 0.3 g·min⁻¹). Total exogenous CHO oxidation over the final (second) hour of running was not modified by MD + F hydrogel (48.25 ± 16.5 g vs. 50.25 ± 16.5 g in MD + F solution) but both were higher than MD + G (41.25 ± 15.0 g). Pettersson et al. (2019) reported exogenous CHO oxidation of 1.22 (0.89–1.66) g·min⁻¹ with MD + F hydrogel, indicating that a higher CHO dose (132 g·hr⁻¹) with hydrogel may result in higher exogenous CHO utilization.

Total rates of CHO and fat oxidation during the steady-state exercise varied between studies, a consequence of different exercise intensities. Therefore, it is of interest to examine relative contributions to total fuel use (Figure 4) as well as absolute rates of oxidation (Table 1). When comparing MD + F in hydrogel versus fluid form, divergent effects were reported. Mears et al. (2020b) and Baur et al. (2019) found no differences in relative substrate contribution during submaximal cycling at 50% Wmax but the nonsignificant increase in fat oxidation (and decrease in CHO oxidation) reported by Baur et al. (2020) is notable, and consistent with Barber et al. (2020). Flood et al. (2020) also reported similar CHO and fat oxidation with CHO and hydrogel ingestion during low-intensity cycling. However, Baur et al. (2019) also noted lower absolute total CHO and fat (to Mears et al. [2020b] in both conditions), despite similar ingestion rates (68 and 78 g·hr⁻¹). McCubbin et al. (2019) did not report differences in substrate oxidation during steady-state running, and neither CHO nor fat oxidation were modified by the hydrogel form during a subsequent incremental test. Barber et al. (2020) however, reported a reduced contribution of endogenous CHO during steady-state running with hydrogel CHO. Glycogen contributed ~60% of total energy expenditure with MD + G and MD + F fluids, but ~50% with MD + F hydrogel. Differences in glycogen use were explained by higher exogenous CHO oxidation in comparison with MD + G fluid, and increased fat oxidation in relation to MD + F fluid (Figure 4). Comparisons between studies employing different exercise modalities for whole-body substrate utilization are difficult as noted previously (Achten et al., 2003).
Table 1  Studies Investigating CHO Hydrogel Formulations With Isocaloric CHO or Placebo Solutions

<table>
<thead>
<tr>
<th>Study</th>
<th>Participants</th>
<th>Design/protocol</th>
<th>Preexercise CHO</th>
<th>Supplement</th>
<th>Performance ∆</th>
<th>Exogenous CHO Ox (g·min⁻¹)</th>
<th>Whole-body substrate Ox (g·min⁻¹)</th>
<th>GI symptoms (hydrogel comparisons only)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baur et al. (2019)</td>
<td>Nine male cyclists</td>
<td>RCT; variable-intensity cycling (55 min SS @ 50% Wmax + 2 × 4 × 2 min @ 80% Wmax + 5 min @ 50% Wmax + 10 × max sprint)</td>
<td>2 hr preexercise</td>
<td>MD + F hydrogel</td>
<td>284 ± 51 W (†2.5%)#</td>
<td>n/a</td>
<td>CHO: 1.50 ± 1.26</td>
<td>No significant Treatment × Time interactions</td>
</tr>
<tr>
<td></td>
<td>(trained, VO₂max = 55.5 ± 3.6 ml·kg⁻¹·min⁻¹)</td>
<td></td>
<td></td>
<td>MD + F solution</td>
<td>281 ± 46 W (†1.08%)#</td>
<td>n/a</td>
<td>CHO: 1.53 ± 1.37</td>
<td>Nonsignificant ↑ fullness vs. MD (ES = 0.54)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MD solution</td>
<td>277 ± 48 W</td>
<td>n/a</td>
<td>CHO: 1.40 ± 1.21</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>n/a</td>
<td>fatty acid oxidation</td>
<td></td>
</tr>
<tr>
<td>McCubbin et al. (2019)</td>
<td>Nine male runners</td>
<td>RCT; fixed-intensity running + TTE (180 min @ 60% VO₂max TTE: SS pace + 2 km·hr⁻¹ ↑ / 3 min)</td>
<td>2 hr preexercise</td>
<td>MD + F hydrogel</td>
<td>744 ± 182 s (†1.6%)#</td>
<td>n/a</td>
<td>CHO: −1.92 ± 0.30</td>
<td>11% vs. 22% incidence of severe symptoms hydrogel vs. MD; F</td>
</tr>
<tr>
<td></td>
<td>(trained, VO₂max = 59.0 ± 8.0 ml·kg⁻¹·min⁻¹)</td>
<td></td>
<td></td>
<td>MD + F solution</td>
<td>90 g·hr⁻¹ in 0.57 L·hr⁻¹</td>
<td>756 ± 187 s</td>
<td>CHO: −1.95 ± 0.30</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MD solution</td>
<td>239 ± 16 W</td>
<td>1.22 ± 0.4</td>
<td>CHO: 2.38 ± 0.25</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>fatty acid oxidation</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pettersson et al. (2019)</td>
<td>12 elite CX skiers: six males, six females (VO₂max =69.1 ± 2.9 and 59.9 ±2.6 ml·kg⁻¹·min⁻¹)</td>
<td>RCT; fixed-intensity CX skiing + TT (120 min @ 70% VO₂max)</td>
<td>1 hr preexercise</td>
<td>MD + F hydrogel</td>
<td>239 ± 16 W</td>
<td>1.22 ± 0.4</td>
<td>CHO: 2.38 ± 0.25</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MD + F solution</td>
<td>132 g·hr⁻¹ in 0.6 L·hr⁻¹</td>
<td>238 ± 16 W</td>
<td>CHO: −0.70 ± 0.06</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>PLA (noncaloric) hydrogel</td>
<td>0</td>
<td>CHO: 2.00 ± 0.26</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>fatty acid oxidation</td>
<td></td>
</tr>
<tr>
<td>Mears et al. (2020b)</td>
<td>Eight cyclists</td>
<td>RCT; fixed-intensity cycling + TT (120 min @ 50% Wmax)</td>
<td>2 hr preexercise</td>
<td>MD + F hydrogel</td>
<td>1,267 ± 102 s</td>
<td>0.9 ± 0.5</td>
<td>CHO: 2.56 ± 0.44</td>
<td>No other significant Treatment × Time interactions</td>
</tr>
<tr>
<td></td>
<td>(well-trained, VO₂max = 62.1 ± 6.9 ml·kg⁻¹·min⁻¹)</td>
<td></td>
<td></td>
<td>MD + F solution</td>
<td>68 g·hr⁻¹ in 0.5 L·hr⁻¹</td>
<td>1,267 ± 102 s</td>
<td>CHO: 2.56 ± 0.44</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>fatty acid oxidation</td>
<td></td>
</tr>
<tr>
<td>Barber et al. (2020)</td>
<td>Nine runners</td>
<td>RCT; fixed-intensity running (120 min @ 60% VO₂max)</td>
<td>Nil: 8 hr fast</td>
<td>MD + F hydrogel</td>
<td>n/a</td>
<td>1.1 ± 0.3</td>
<td>CHO: 2.60 ± 0.75</td>
<td>No significant Treatment × Time interactions</td>
</tr>
<tr>
<td></td>
<td>(well-trained, VO₂max = 63 ± 3.6 ml·kg⁻¹·min⁻¹)</td>
<td></td>
<td></td>
<td>MD + F solution</td>
<td>1.1 ± 0.3</td>
<td></td>
<td>fatty acid oxidation</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>MD + G solution</td>
<td>0.9 ± 0.5</td>
<td></td>
<td></td>
<td>fatty acid oxidation</td>
</tr>
</tbody>
</table>

(continued)
Table 1  (continued)

<table>
<thead>
<tr>
<th>Study</th>
<th>Participants</th>
<th>Design/protocol</th>
<th>Preexercise CHO</th>
<th>Supplement</th>
<th>Performance Δ</th>
<th>Exogenous CHO Ox (g·min⁻¹)</th>
<th>Whole-body substrate Ox (g·min⁻¹)</th>
<th>GI symptoms (hydrogel comparison only)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flood et al. (2020)</td>
<td>14 cyclists: seven males, seven females (recreational/trained, VO₂max = 56.4 ± 7.6 ml·kg⁻¹·min⁻¹; 54.3 ± 12.3 ml·kg⁻¹·min⁻¹)</td>
<td>RCT; fixed-intensity cycling + TT (90 min @ 45% VO₂max)</td>
<td>3 hr preexercise</td>
<td>MD + F hydrogel</td>
<td>192 W (↑14%)</td>
<td>CHO: 1.73 ± 0.75</td>
<td>IFABP (end exercise)</td>
<td>▲ Water (~500 pg·ml⁻¹ vs. hydrogel (~150 pg·ml⁻¹))</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Self-selected “CHO rich”</td>
<td>MD + F solution</td>
<td>190 W (↑13%)</td>
<td>FAT: 0.21 ± 0.35</td>
<td>MD + F [by ~70 pg·ml⁻¹]</td>
<td>Lactulose: Rhamnose ↓ in MD + F and hydrogel**</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>PLA 90 g·h⁻¹ in 0.78 L·h⁻¹</td>
<td>168 W</td>
<td>n/a</td>
<td>CHO: 1.35 ± 0.40</td>
<td>No significant treatment effects for GI symptoms. Fullness ↑ in MD + F and hydrogel**</td>
</tr>
</tbody>
</table>

Note. RCT = randomized crossover trial; SS = steady state; TTE = time to exhaustion; TT = time trial; CHO = carbohydrate; ER = energy requirement; CX = cross-country (skiing); IFABP = Intestinal fatty acid binding protein; MD = maltodextrin; F = fructose; G = glucose; GI = gastrointestinal; PLA = noncaloric placebo. *Performance change not significantly different (p > .05). Performance changes in Baur et al. (2019) are relative to MD condition and in Flood et al. (2020) relative to water. **Gastrointestinal effects significantly different (p < .05) to placebo.
Gastrointestinal discomfort was reported in all studies with similar responses from the hydrogel trial. McCubbin et al. (2019) reported a slightly, but not significantly, higher overall incidence of GI symptoms while running with MD + F fluids than MD + F hydrogel (22% vs. 11%). In cycling, Mears et al. (2020b) and Baur et al. (2019) reported slightly increased stomach fullness with CHO hydrogel ingestion but similarly, differences were not significant. Fullness in Flood et al. (2020) did not differ between CHO hydrogel and MD + F. Baur et al (2019) also reported moderately, but not significantly increased nausea with the hydrogel compared with MD + F fluids (ES = 0.53, p = .23). Data from Pettersson et al. (2019) reported low levels of GI issues during skiing in cold conditions in either trial, but a small, nonsignificant decrease in stomach rumbling with the hydrogel compared with MD + F fluids (ES = 0.53, p = .23). Data from Pettersson et al. (2019) reported low levels of GI issues during skiing in cold conditions in either trial, but a small, nonsignificant decrease in stomach rumbling with the hydrogel compared with MD + F. Baur et al (2019) also reported moderately, but not significantly increased nausea with the hydrogel compared with MD + F fluids (ES = 0.53, p = .23).

Discussion

This is a timely investigation of the evidence that intake of hydrogel encapsulations of multiple transportable CHO during prolonged endurance exercise provides benefits over traditional sports drinks, in response to the recent interest in newly available commercial hydrogel products. We summarized studies where hydrogels, formed by combining MD + F with gelling agents such as alginate and pectin, were compared with typical CHO fluids containing single or multiple CHO sources, or a noncaloric hydrogel treatment. Despite marketing claims and lay media discussion about MD + F products in hydrogel format, currently available studies fail to show benefits in terms of muscle oxidation of exogenous CHO, GI comfort, or performance. The current literature, comprised of robust randomized controlled trials, is small and includes nuances around total substrate oxidation and gut comfort due to exercise mode and intensity, as well as total CHO intake. Furthermore, the conditions under which it is promoted to achieve its key benefits (high rates of CHO intake during prolonged exercise at high absolute and relative intensities) have not been investigated, potentially due to the challenge of involving elite competitors within traditional research protocols and the technical challenges of undertaking measurements of interest (e.g., gastric emptying, tracer determined substrate oxidation) under these conditions. While further studies with relevant protocols are needed to investigate the putative benefits of these products, the present literature fails to endorse the marketing claims.

Over the past 5 years, CHO-containing drinks that achieve hydrogel encapsulation within the gut have become commercially available, with claims that they achieve superior CHO delivery to the muscle for lower GI distress, leading to performance benefits over traditional CHO-containing sports products (Maurten, 2020; Sutehall et al., 2018). CHO hydrogel products have attempted to improve on existing nutrition recommendations (use of CHO with multiple transporters [Jeukendrup, 2010] and gut training [Jeukendrup, 2017]) to target the need for high-performance athletes to achieve high CHO availability during prolonged events conducted at high relative and absolute intensities (Burke et al., 2019). Although there is evidence that these products achieve gelation within the acidic stomach environment as claimed (Marciani et al., 2019; McCubbin et al., 2019), subsequent effects on gastric emptying, intestinal absorption, and delivery to the muscle during exercise remain largely untested. Our review summarized the available literature on the use of these products during prolonged exercise in terms of GI comfort, substrate utilization, and performance. The scarcity and heterogeneity of study protocols prevents a meta-analytical approach from producing meaningful results. However, in view of community interest and marketing claims, we felt it was timely to collate the findings of available studies in narrative form to summarize the overall findings and alert researchers to the need for particular protocols.
The current literature involves studies of subelite athletes (although one study of elite performers is noted [Pettersson et al., 2019]), exercising over 2–3 hr at intensities of approximately 45–70% VO$_2$peak. Such protocols likely represent pragmatic choices based on subject availability and the technical requirements around the steady-state conditions needed to monitor muscle substrate use during exercise (Robert et al., 1987). According to our review, although the GI handling of these products has not been directly measured, these studies have failed to find evidence of increased oxidation of exogenous CHO when MD + F is provided in hydrogel form compared with conventional solutions. Indeed, rates of exogenous CHO oxidation derived from $^{13}$C isotopic tracer techniques are in line with results of previous studies of traditional CHO-containing fluids (King et al., 2019; O’Brien et al., 2013), or in the case of studies in which direct comparison has been made between CHO-matched fluids and hydrogel products, no differences in exogenous CHO use has been detected (Barber et al., 2020). Since the rate-limiting step of exogenous CHO oxidation is believed to lie in the GI tract and not cellular glucose uptake (Hawley et al., 1994), this indirectly indicates that CHO hydrogels have not achieved a net benefit to the gastric emptying/intestinal CHO transport processes.

The studies published to date confirm previous knowledge that whole-body CHO utilization is altered by the intake of CHO during exercise (Pettersson et al., 2019) and by the choice of multiple transportable CHO sources that increase the capacity for total intestinal absorption (Barber et al., 2020; Baur et al., 2019). In general, Figure 4 suggests that hydrogel encapsulation per se does not change total CHO oxidation during exercise (Baur et al., 2019; Flood et al., 2020; McCubbin et al., 2019; Mears et al., 2020b). However, Barber et al. (2020) reported a decrease in endogenous CHO utilization and increase in fat oxidation when high rates of CHO intake were consumed in hydrogel form versus fluid. Muscle glycogen sparing effects have been reported in several studies of CHO intake during endurance exercise, but performance improvements have not been consistently observed (Newell et al., 2014). Previous work from our group (King et al., 2018) and others (Smith et al., 2010) has shown that the muscle glycogen response to CHO ingestion is dose dependent. Precise mechanisms to explain this effect have not been investigated, but likely sit within the cellular flux through the glycolytic pathway and the interaction of exogenous glucose and glucosyl units liberated from glycogen at glucose-6-phosphate. Exogenous CHO more consistently reduces liver glycogen use as long as the ingested dose is sufficient to inhibit hepatic glycogenolysis and glucose output (Gonzalez & Betts, 2019). However, liver glycogen capacity is much smaller than muscle (~100 g vs. ~400–500 g) and complete liver glycogen sparing during exercise has only been reported with an extremely high CHO dose (Jeukendrup et al., 1999). The ingested doses in the reviewed studies cover a wide range, and only one study estimated whole-body endogenous glycogen utilization using expired $^{13}$CO$_2$ tracer methods, which do not account for specific liver and muscle contributions; conclusions around a dose effect with CHO hydrogels are not possible at this stage. However, if an event requires maximized CHO availability, recommendations to saturate intestinal CHO transporters should remain if athletes tolerate these doses in terms of GI distress.

As has been the case in previous investigations of CHO feeding during exercise, methodological differences between studies do not allow firm interstudy comparisons. Factors such as CHO

![Figure 4](https://example.com/image.png)  
**Figure 4** — Comparison of relative contributions to energy expenditure from fat (black bars) and carbohydrate (white bars) with maltodextrin and fructose (MD + F) ingestion in fluid and hydrogel form. Exogenous (dotted bars) and endogenous (hashed bars) contributions shown where data were available. MD + F = maltodextrin + fructose; CHO = carbohydrate.
dose, as well as timing, exercise intensity, mode and duration, prefeeding state, and potentially training status create differences in outcomes. Although such variability creates difficulty in piecing together an emerging literature, it is important to recognize that many of these factors may be fixed or characteristic of specific sporting events. Further studies, particularly those using multiple conditions within the same investigation, might help to isolate any conditions under which hydrogel CHO might provide an advantage. Exercise intensity has important implications for intestinal absorption due to the diversion of splanchnic blood flow at higher intensities (ter Steege & Kolkman, 2012). Therefore, potential mechanistic benefits of CHO hydrogels in terms of GI tolerance and absorption may only be observed in the higher intensity domains, presenting an opportunity for future research. To date, only one study (Pettersson et al., 2019) has investigated what may be considered higher intensity exercise. Running also utilizes less localized muscle recruitment compared with cycling, resulting in increased surface area blood flow and a greater reduction of splanchnic blood flow (de Oliveira et al., 2014). Similar responses are likely in cross-country skiing given the whole-body nature of the sport and may even demand higher CHO utilization (Losnegard et al., 2014). While body mass is not considered a methodological factor for exogenous CHO oxidation (Jeukendrup, 2010), the combination of differences in exercise mode and intensity in the reviewed studies does not allow for firm extrapolations to higher intensity exercise, which may also benefit further from optimal, that is, at intestinal saturation, CHO dosing. Due to the increased duration of training and competition, cyclists may be at an advantage over runners if habituated to CHO intake. Increased CHO exposure causes a “gut training” effect (Cox et al., 2010), leading to enhanced GI tolerance of ingested CHO during exercise. Gut tolerance is therefore a further methodological factor that differentiates between exercise modality and may mean potential GI benefits to CHO hydrogels may be more likely in individuals with lower natural CHO tolerance or with less habitual CHO gut training. This would serve as a useful consideration or screening tool in future study design.

Typically, the intake of solutions with high CHO content during prolonged exercise delays gastric emptying and is associated with higher incidence of GI distress (Rehrer et al., 1992). A positive finding from this review is that MD + F hydrogel formulations were generally well tolerated across the range of doses and exercise protocols that were examined. However, hydrogel solutions did not improve GI tolerance per se above comparable CHO sources in traditional fluid form. It remains to be seen if they systematically reduce GI symptoms at higher doses approaching and above intestinal saturation, due to specific interaction with the digestive system. Reports of a slight increase in gastric fullness associated with the hydrogel are of interest (Baur et al., 2019), since even if there is a subsequent increase in gastric emptying associated with the formation of the gel, it may create an initial sensation of fullness. Quantitative measures of GI barrier function and damage reported by Flood et al. (2020) evidence that CHO hydrogels do not provide a further protective effect to the intestinal membrane over typical CHO ingestion. These data do however, confirm the preventative role of CHO for enterocyte injury and small intestine permeability during endurance exercise (Snipe et al., 2017).

Although an enhancement of gastric emptying is the mechanism most heavily marketed in support of the use of the hydrogel sports drinks, the importance of gastric emptying in the whole process of delivering CHO from the mouth to the muscle mitochondria, and any benefits achieved by hydrogels, are difficult to ascertain. So far, the available evidence is limited to a report by Sutehall et al. (2020) that a commercially available MD + F hydrogel increased gastric emptying compared with G + F and MD + F solutions when consumed as a bolus at rest, with time to empty half the drink being nearly twice as fast for MD + F hydrogel (21 min) than its fluid counterpart (37 min). Despite significantly less volume remaining in the stomach at 30 min, differences became increasingly smaller thereafter (Sutehall et al., 2020). Since none of the currently available studies have attempted to directly measure gastric emptying during exercise, it is not possible to comment on what occurs under these circumstances. However, according to the general literature, gastric emptying is not considered to be the rate-limiting step in determining the availability of exogenous fuels consumed during exercise (see review by Rowlands et al. (2015)) and its measurement includes artifacts and practical difficulties. Furthermore, without using complex invasive measures (Shi et al., 1995), intestinal absorption cannot be directly measured; therefore, the endpoint of muscle CHO oxidation is used to reflect the contribution of a combination of gut processes between consumption and delivery to the mitochondria. The evidence collected from the currently available studies does not show clear evidence of differences in the overall process.

Even if hydrogel-encapsulated CHO can be shown to have different gut characteristics per se, the effects of the amount, timing, and pattern of intake of CHO sources on gastric emptying are among the many interacting factors that should be considered in the current story. This is of interest since in many sporting events, the pattern of intake is dictated by the availability of fluids at breaks or feed stations rather than continuous or spontaneous access. Recent work by Mears et al. (2020a) reported that mean and peak exogenous CHO oxidation is altered slightly (but significantly) by CHO timing, with better outcomes associated with consuming sources every 20 min compared with 5-min intervals. Meanwhile, Menzies et al. (2020) reported better endurance when CHO intake commenced early during a running protocol than later delivery. Therefore, future research should systematically investigate conditions under which a hydrogel CHO might be used. Although this will involve a large number of permutations of characteristics, we note in particular, that scenarios from which current testimonials about the use of hydrogels have emanated have not been investigated. These include use by elite athletes in events such as the marathon requiring high CHO availability from endogenous and exogenous sources to fuel prolonged exercise (~2 hr+) of high relative and absolute intensities (Maurten, 2020; Sutehall et al., 2018). Here, the combination of issues such as high fuel requirement, high risk of gut distress, and a practical requirement for small fluid volumes intersect, making them a priority for study.

In conclusion, a small number of studies have investigated the use of commercially available CHO hydrogels to deliver CHO during exercise. So far, data do not support the claimed benefits of enhanced CHO delivery to the muscle, reduced GI distress, and better performance compared with the use of traditional CHO solutions. Future research should focus on dose and timing of hydrogel ingestion, higher exercise intensities where GI issues are more prevalent and CHO absorption more greatly impaired, and further mechanistic insights around endogenous CHO responses. Athletes may continue to use CHO hydrogels to meet current guidelines for endurance nutrition practices if this is their preference. While no disadvantages around the use of these specialized sports products appear to be present, based on current evidence, they do not confer metabolic or performance advantages over typical CHO ingestion strategies.
Acknowledgments

All authors contributed to the preparation of this manuscript. The authors declare no conflicts of interest in the preparation of this review.

References


