Microbial metabolism of transparent exopolymer particles during the summer months along a eutrophic estuary system

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This study explores the role of transparent exopolymer particles (TEP) as an additional carbon source for heterotrophic microbial activity in the eutrophic Qishon estuary. From the coastal station and upstream the estuary; TEP concentrations, β-glucosidase activity, bacterial production and abundance have gradually increased. TEP were often found as bio-aggregates, scaffolding algae, detritus matter and bacteria that likely formed “hotspots” for enhanced microbial activity. To further demonstrate the link between TEP and heterotrophic bacterial activity, confined incubations with ambient and polysaccharide-enriched estuary water were carried out. Following polysaccharide addition, elevated (~50%) β-glucosidase activity rates were observed, leading to TEP hydrolysis. This newly formed bioavailable carbon resulted in significantly higher growth rates, with up to a 5-fold increase in heterotrophic bacterial biomass, comprising mostly high nucleic acid content bacteria. Taking together the findings from this research, we conclude that even in highly eutrophic environments heterotrophic bacteria may still be carbon limited. Further, TEP as a polysaccharide matrix can act as a metabolic surrogate, adding fresh bioavailable carbon through tight associations with bacteria in eutrophic ecosystems such as the Qishon estuary.

Keywords: heterotrophic bacteria, TEP, estuary, ectoenzyme, β-glucosidase, polysaccharides, BGE, BCD

Introduction

Transparent exopolymer particles (TEP) are planktonic, acidic-polysaccharide hydrogels that are ubiquitous in various marine and freshwater environments (Passow, 2002; Bar-Zeev et al., 2015). TEP are clear and therefore only detected by specific stains such as alcian blue (Allredge et al., 1993; Decho and Lopez, 1993). In the last two decades, TEP were shown to mediate diverse biochemical cycles in aquatic environments; TEP promote aggregate formation and sedimentation (Passow et al., 2001; Simon et al., 2002; Engel, 2004) by providing a scaffold for marine or lake “snow” (Mari and Kirboe, 1996; Grossart et al., 1997, 2006; Berman and Viner-mozzini, 2001). TEP may also serve as surface and substrate for microbes, forming “hotspots” of intense microbial activity (Azam, 1998; Simon et al., 2002; Azam and Malfatti, 2007), thus fueling the microbial loop and impacting food web structure in various ecosystems (Simon et al., 2002; Beauvais et al., 2006; de La Rocha and Passow, 2007).
Estuary systems are dynamic, semi-enclosed coastal environments where open seawater is partly diluted by fresh river water (Pritchard, 1967; Elliott and McLusky, 2002). Most estuaries worldwide are turbid and highly productive due to constant allochthonous nutrient inputs (Pinckney et al., 2001). Convergence of estuaries and open seawater often result in some of the steepest physiochemical gradients, including light penetration and oxygen concentration (Malone et al., 1996; Hall and Pearl, 2011).

The highly stratified Qishon estuary at the northern coast of Israel flows into the southeastern Mediterranean Sea, producing a steep eutrophic to oligotrophic gradient (Herut et al., 1999; Eliani-Russak et al., 2013). In the summer season, the high air temperature, low flows (0.02–0.2 m$^3$ s$^{-1}$) (Vachtman et al., 2013) and high nutrient load (as nitrate, $\sim$240 tons per year) produce a stratified, nutrient-rich 7 km long estuary (Eliani-Russak et al., 2013). The estuary bottom layer is constantly saline (34.1–39.9) and exhibit high nutrient concentrations (1–85 $\mu$M NO$_3$ and 0–9.3 $\mu$M PO$_4$) throughout a spatial gradient from inland and toward the coast (Eliani-Russak et al., 2013).

In estuary ecosystems, TEP were found to play a significant role in organic matter cycling through degradation, aggregation and sedimentation processes (Simon et al., 2002; Barrera-Alba et al., 2009; Wetz et al., 2009; Mari et al., 2012; Sun et al., 2012). Hydrodynamic conditions, governed mainly by winds and tides, were reported to play a key role in TEP cycling between estuary and coastal environments (Mari et al., 2012; Sun et al., 2012). Fluvial cations passing through the estuary may also alter TEP formation (Simon et al., 2002; Wetz et al., 2009; Sun et al., 2012). In addition to its physicochemical roles in estuary systems, TEP were also suggested to act as vector for different types of pathogens (Lyons et al., 2007; Wetz et al., 2009).

In this study, we focus mainly on the metabolic feedback loops between TEP and the heterotrophic microbial community at a eutrophic estuary system. To do so, we sampled three stations along the Qishon estuary system, while conducting confined-laboratory incubation experiments. Our results shed new light on carbon limitation and polysaccharide metabolism by heterotrophic bacteria, highlighting the importance of TEP as a carbon source in a highly eutrophic environment.

**Materials and Methods**

**Sampling Strategies and Bioassay Designs**

Four stations were sampled; three along the Qishon estuary (E1; Maagan, E2; Yulius and E3; Histadrut) and one reference coastal station (Sh) (*Figure 1, Table 1*) during the summer months (September 2013 and June to August 2014). Temperature, salinity, dissolved oxygen (DO) and turbidity were measured in each sampling site throughout the water column using an YSI (Model 6600) profiler system. Near surface (~20 cm deep) and bottom water samples were collected in each sampling location using a 3.1 L Niskin bottle. Water samples were analyzed for TEP, bacterial production and abundance, as well as $\beta$-glucosidase activity, dissolved inorganic nutrients and chlorophyll a (Chl a).

To elucidate the role of TEP as a labile carbon source for heterotrophic bacteria, we conducted two bottle incubation experiments simulating high TEP scenarios during June and August 2014. The incubation bottles were pre-washed with 10% hydrochloric acid, rinsed three times with Milli-Q water followed by three times with ambient water. At each sampling date, bottles were filled with water samples (1 L) collected from the surface and bottom of the three estuary stations ($n = 9$). Samples were then pre-filtered (1 $\mu$m filter, PALL co.) to remove most of the particulate matter, including large TEP, micro algae, filamentous cyanobacteria and grazers, while retaining planktonic bacteria. The pre-filtered water was then supplemented with gum-xanthan (GX, Sigma G1253); a commercial pure polysaccharide secreted by the bacterium *Xanthomonas campestris* (Passow and Alldredge, 1995). Gum-xanthan additions were prepared by dissolving 5 mg of GX in deionized water (100 mL) and homogenized (Thomas Scientific Model D1000) for 15 min to form concentrated small-suspended particles (Rahav et al., 2013). GX additions yielded TEP concentrations ranging from 1532 to 1648 $\mu$g GX L$^{-1}$, which were ~8-fold higher relative to the pre-filtered, un-amended controls. The pre-filtered control and GX supplemented bottles were then incubated for 2 days under ambient temperature (25°C) in complete darkness to minimize primary production through photosynthesis. Subsamples were collected and analyzed for bacterial production, respiration and abundance, $\beta$-glucosidase activity, TEP and Chl a at the beginning (T$\text{_0}$) and at the end (T$\text{_2}$d) of the microcosm experiment. Bottles were thoroughly shaken before collecting the subsamples to resuspend any biofouling/fouling that may developed on the polycarbonate walls and aggregates that could have sedimented during the incubations.

**Dissolved Inorganic Nutrients**

Water samples were collected in 15 mL acid-washed plastic scintillation vials. Nutrient concentrations were determined using a segmented flow Seal Analytical AA-3 system (Krom et al., 1991; Kress and Herut, 2001). The limits of quantitation...
of nitrate + nitrite, ammonium and phosphorus measurements were 0.19 µM, 0.15 µM and 0.02 µM, respectively. The limits of detection (twice the standard deviation of the blank) were 0.08 µM for nitrate + nitrite, 0.06 µM for ammonium and 0.008 µM for phosphorus.

**Chlorophyll a (chl a) Biomass as Algal Proxy**

Water samples (150 mL) were passed through 0.7 µm glass fiber filters (Whatman, Lot: 1825025) and stored at −20°C, protected from light. Samples were extracted in 5 mL of 90% acetone overnight at 4°C in the dark. Chl a concentrations were determined using a luminescence Trilogy® fluorometer (7200-000) with a 436 nm excitation filter and a 680 nm emission filter (Holm-Hansen et al., 1965).

**Heterotrophic Bacterial Abundance (BA) and Specific Growth Rates**

Water samples (1.8 mL) were fixed with 6 µL of 50% glutaraldehyde (Sigma, G7651), incubated in room temperature for 10 min, flash-frozen in liquid nitrogen and stored at −80°C until further analyses. Prior to counting, the samples were thawed at 37°C for 2 min, stained with 1:105 vol:vol nucleic acid SYTO®9 (Life technologies S-7580) in the dark for 10 min and analyzed by an Attune® acoustic focusing flow cytometer (Applied Biosystems). Samples were excited with Argon lasers (488 and 405 nm) with a flow rate of 25 µL min⁻¹. Heterotrophic bacteria were specifically enumerated by subtracting autotrophic bacteria (detected by Chlorophyll a auto-fluorescence) from the total bacterial abundance. One-micron beads (Polysciences) were used as size calibration standard. Heterotrophic bacterial abundance was converted to carbon biomass using a factor of 20 fg C per heterotrophic bacterial cell (Lee and Fuhrman, 1987). High and low nucleic acid content bacteria (HNA and LNA respectively) were differentiated by side scattering (SSC) and SYTO 9 fluorescence (488 nm excitation) that was detected by a 530/30 nm band pass filter.

Heterotrophic bacterial (HB) growth rates (k) were calculated according to the following equation:

$$k_{day} = \frac{ln(BA_{T2}/BA_{T0})}{T_2 - T_0}$$  \hspace{1cm} (1)

To calculate k, heterotrophic BA was enumerated at the begging (T₀) and the end (T₂) of the bottle incubations.

**Bacterial Production Rates (BP)**

BP was measured using the [4,5-³H]-leucine incorporation method (Simon et al., 1990). Briefly, three aliquots (1.7 mL each) from each sample were incubated with 100 nmol L⁻¹ of [4,5-³H]-leucine (Perkin Elmer USA, Lot: 1804258) for 4 h in room temperature in the dark. Triplicate trichloroacetic acid (TCA) inactivated samples served as controls. The incubations were terminated with 100 µL of cold (4°C) TCA (100%), followed by centrifugation (Smith and Azam, 1992) to form a pellet. After removing the supernatant and adding 1 mL of scintillation cocktail (Ultima-Gold, Quick-Safe A) to each tube, the samples were counted using a TRI-CARB 2100 TR (Packard TRI-CARB 2100 TR) liquid scintillation counter. Leucine incorporation was converted into carbon assimilation by a conservative factor of 3.1 kg C mol⁻¹ with an isotope dilution factor of 2.0 (Simon and Azam, 1989).

**Dissolved Oxygen (DO) and Bacterial Respiration Rates (BR)**

Oxygen was measured using the Winkler titration procedure (Carpenter, 1965). Briefly, water was immediately fixed with MnSO₄ and KI + NaOH and sealed without headspace in 300 mL Winkler bottles (Wheaton® 227497-11). H₂SO₄ was later added, and samples were titrated with Na₂S₂O₃ using a
Metrohm 785 DMP titrino auto-burette and double platinum electrode (end-point titration precision, ± 1 µmol L⁻¹) similarly to Kress et al. (2014).

Respiration rates were determined by the following equation:

\[ \text{Respiration} = \frac{\text{DO}_{(T0 \text{ dark})} - \text{DO}_{(T48 \text{ dark})}}{48 \text{ h}} \]  

(2)

Where \(\text{DO}_{(T0 \text{ dark})}\) is the initial dissolved oxygen concentration and \(\text{DO}_{(T48 \text{ dark})}\) is the dissolved oxygen concentration after 48 h incubation. We assumed that in all bottle incubations bacterial respiration (BR) accounted for ∼90% of the dark respiration of the entire microbial community (see result and discussion for further details).

**Bacterial Carbon Demand (BCD) and Bacterial Growth Efficiency (BGE)**

BCD was defined as the sum of carbon assimilation measured by bacterial production (BP) and carbon oxidation determined through heterotrophic microbial respiration (BR). Oxygen respiration was converted into carbon consumption assuming a respiratory quotient (RQ) of 1 (del Giorgio and Cole, 1998; Anesio et al., 2003; Smith and Prairie, 2004):

\[ \text{BCD} = \text{BP} + \text{BR} \]  

(3)

BGE was calculated as follows:

\[ \text{BGE} (\%) = \frac{\text{BP}}{\text{BP} + \text{BR}} \times 100 \]  

(4)

**TEP Concentrations and Visualization with Associated Bacteria**

Water samples (100 mL) were gently (<150 mbar) filtered through a 0.4 µm polycarbonate filter (GE Water & Process Technologies). Filters were then stained with a solution of 0.02% alcian blue (AB), 0.06% acetic acid (pH of 2.5), and the excess dye was removed by a quick deionized water rinse. Filters were then immersed in sulfuric acid (80%) for 2 h, and the absorbance (787 nm) was measured spectrophotometrically (Thermo GENESYTM). AB dye was calibrated using GX as a purified polysaccharide (Passow and Alldredge, 1995). A factor of 0.74 was used to convert from GX equivalents to carbon (Engel and Passow, 2001). This conversion factor was used as proxy, since TEP chemical composition is likely to change between different marine and fresh water environments.

To visualize TEP with bacterial associations, samples (100 mL) were filtered gently (<150 mbar) onto 0.4 µm polycarbonate filters (GE Water and Process Technologies) and stained with 0.2 µm pre-filtered AB solution and 30 µL SYTO®9 (250 µg mL⁻¹) for TEP and bacterial identification, respectively (Bar-Zeev et al., 2011). After 7 min incubation in the dark, filters were washed with deionized water (5 mL) and mounted on a coated Cytoclear (Clearing Slides, GE Osmonics Labstore) slide. TEP images were taken with bright-field (Nomarski) illumination; bacteria were visualized with an epifluorescence microscope (Olympus BX50 microscope) equipped with FITC filter (excitation: 488 nm; emission: 520 nm). The detection of Chl a fluorescence was examined by a specified filter set (ex: 450 nm, em: 680 nm).

\[ \beta - \text{Glucosidase Activity Rates (\(\beta\)-glu)} \]

The hydrolytic activity of β-glucosidase was determined by cleavage rates analysis of a conjugated fluorogenic substrate, 4-methylumbelliferyl (MUF)-β-D-glucopyranoside (Sigma M3633) (Hoppe, 1993; Luna et al., 2012). Briefly, the substrate (final concentration of 50 µM) was added to a 1 mL water sample in triplicate and incubated in the dark at ambient temperature (25°C) for 24 h. The increase in fluorescence was measured at 365 nm excitation and 455 nm emission (GloMax®-Multi Detection System E9032). Reads were normalized against a calibration curve ranging from 0 to 250 µM MUF (\(R^2 = 0.99\)). We used a conversion factor of 72 to convert the hydrolyzed conjugated fluorogenic substrate, measured as nM MUF to µg carbon (Hoppe, 1993).

**Statistical Analysis**

Data is displayed as average with error bars signifying one standard deviation (\(n = 7–9\)). All field relationships were determined with a Pearson correlation test (\(n = 7–9\) and \(P < 0.01\)). Throughout all bottle incubations, significant differences between control and enriched microcosms were determined using Student’s t-test with paired two-tailed distribution (\(P < 0.01\)). Changes in bacterial abundance were evaluated using One-Way analysis of variance (ANOVA) followed by Fisher LSD multiple comparison post-hoc test with a confidence of 95% using the XLSTAT software.

**Results and Discussion**

**Physicochemical Properties of the Qishon Estuary during the Summer Season**

The Qishon estuary is a 7 km long and shallow (~3 m deep) stream that flows through an industrial environment. During our study (June–September), no precipitation events were recorded (www.ims.gov.il), resulting in minimal freshwater supplements and slow flow rates (0.02–0.2 m s⁻¹) (Vachtman et al., 2013). During the summer, daytime solar irradiance was high (~1500 µmol quanta m⁻² s⁻¹); yet, the turbid nature of this estuary system minimized light penetration to surface layers (~0.2 m), limiting phytoplankton photo-damage (MacIntyre and Cullen, 1996). The surface of the estuary (upper 0.2 m) was saline (18–29), whereas the bottom (~3 m) was significantly saltier ranging from 36 to 39 (Figure 2, Table 1). The high salinity at the bottom was mainly due to eastern Mediterranean seawater percolating through from the sediment (Herut and Kress, 1997; Eliani-Russak et al., 2013). At all locations and sampling periods, temperature was mostly uniform (25–27°C) throughout the water column (Figure 2, Table 1). However, due to profound salinity differences, the estuary bottom layer was much denser than the surface (14 and 7.5 kg m⁻³ respectively), resulting in a strong stratification. The stable water column has possibly restricted vertical mixing and resulted in oxygen-depleted, low pH water at the bottom (Figure 2, Table 1) due to high microbial metabolic activities.

Effluents of two fertilizers plants, an oil refinery and a sewage treatment plant often flow into the Qishon estuary system, resulting in a eutrophic gradient (Table 1). Station E3, is located
upstream and close to the above industrial effluents. As a result, dissolved inorganic nitrogen (NO₂ + NO₃, NH₄, hereafter DIN) and phosphorus (DIP) concentrations were exceptionally high (543 ± 623 µM and 8.8 ± 4.5 µM respectively), with low DO levels (2.3 ± 2.2 mg L⁻¹) indicating high bacterial activity (Table 1). Downstream, station E2 exhibited lower inorganic nutrient concentrations than E3, possibly due to gradual dilution, re-mineralization and sedimentation of aggregates (Table 1). Station E1 is located at the mouth of the estuary in a fishing harbor, and is therefore affected by small vessels, as well as mixing between the Qishon stream at the surface and coastal water at the bottom. Therefore, DIN was high at the surface (515 µM) and decreased by 7-fold near the bottom (72 µM). DO levels varied from 6.6 mg L⁻¹ at the surface to 3.3 mg L⁻¹ at the bottom and similar to E3 indicate high heterotrophic metabolism. Along the entire estuary, turbidity was significantly higher at the bottom (29 ± 11 NTU) than in the surface (8 ± 3 NTU) layer.

At the same time, a coastal station (Sh) uninfluenced by the estuary (Figure 1) was monitored as reference. The coastal station exhibited typical oligotrophic southeastern Mediterranean characteristics; low turbidity (~1.5 NTU), warm temperature (25 ± 3°C), high salinity (39 ± 0.4) and well oxidized water column (Figure 2, Table 1). DIN and DIP concentrations were 2–3 orders of magnitude lower than in the Qishon estuary (Table 1).

The eutrophic conditions in the Qishon water were probably due to anthropogenic effluents and the slow flow rate during the dry summer, resulting in a sharp nutrient gradient from the estuary toward the Mediterranean coast.

### Linking Phytoplankton and Bacterioplankton to TEP along the Qishon Estuary System

Phytoplankton biomass measured as Chl a, TEP concentrations, and heterotrophic bacterial abundance were all significantly higher in the Qishon estuary system compared with the reference coastal station (Figure 3). Chl a concentrations were 40–100-folds higher in the estuary than at the coastal station (Sh). A positive correlation was also found between phytoplankton concentration and distance from the coastal environment ($R^2 = 0.92, P < 0.001, n = 7$), peaking at station E3; 24 ± 1 µg Chl a L⁻¹ (Figure 3A and Figure S1A). We suggest that this significant relationship point on intense eutrophication, which resulted from sharp increase in nutrient availability (Table 1). However, phytoplankton was equally distributed along the depth
FIGURE 3 | Parameters sampled from the surface and bottom layers, along the eutrophic gradient; from Qishon estuary system (E3 to E1) to the coastal reference station (Sh). Water samples were analyzed for: Chlorophyll a as phytoplankton proxy (A), TEP concentrations (B), Microscopic visualization of a representative bio-aggregate (C), heterotrophic bacterial abundance (D), bacterial production (E), and β-glucosidase activity (F). Bars represent the average of three independent sampling dates with corresponding standard deviation. To visualize bio-aggregates water were sampled from a representative station (E1). Bacterial clusters were stained with SYTO 9 (bright green) and captured with epifluorescent microscopy. Bright field microscopy was used to visualize transparent exopolymers particles (TEP) stained with alcian blue (as light blue), and algal cells (as dark green). Images were superimposed using Image J software (http://rsbweb.nih.gov). Chlorophyll a pigment was not detected by autofluorescence within the bacterial clusters.

profile of most measured stations, despite the turbid conditions in the Qishon estuary (Figure 3A, Table 1).

Along the Qishon estuary, TEP concentrations ranged between 225 and 1059 µg GX L⁻¹ and were higher than the coastal station by 2–5-fold (Figure 3B). TEP were also found to increase linearly with the eutrophic gradient ($R^2 = 0.83, P < 0.001, n = 7$), peaking at station E3 (Figure 3B and Figure S1B). However, TEP normalized to phytoplankton concentrations (µg GX Chl a⁻¹ L⁻¹) followed an opposite trend and were up to 21-fold higher at the coastal station than the estuary. This opposing trend suggest that in the eutrophic Qishon estuary high TEP concentrations were the result of high phytoplankton biomass, while in the oligotrophic coastal environment induced physiological stress due to nutrient limitation may result in higher polysaccharide excretion per cell (Berman-Frank et al., 2007; Bar-Zeev et al., 2011, 2013). Moreover, it is possible that the high microplankton biomass within the estuary promoted viral infection that may result in cell lysis and TEP formation (Vardi et al., 2012; Lønborg et al., 2013).

TEP concentrations were consistently greater at the bottom in all estuary stations (Figure 3B). Sampled TEP were visualized as tightly packed, micrometer-size (hundreds of µm) bio-aggregates with numerous bacterial clusters held in a mixture of micro-algal cells (Figure 3C). No Chl a pigment was detected in the bacterial clusters based on epifluorescence microscopy, indicating that these microbial communities consisted primarily of heterotrophs (Figure 3C). These findings highlight the direct association between TEP mediated-aggregates and heterotrophic bacterial clusters. While drifting as aggregates, these newly formed, fresh-TEP are more labile (Passow, 2002) and are therefore likely to be utilized as substrates for bacterial growth. We suggest that the higher TEP concentrations at the bottom of the estuary were possibly due to polysaccharide release by benthic suspension feeders (McKee et al., 2005; Heinonen et al., 2007) and sedimentation processes, as previously shown...
for other estuaries and aquatic environments (Logan et al., 1995; Passow et al., 2001; Simon et al., 2002; Beauvais et al., 2006).

Heterotrophic bacterial abundances (BA) were 4–30-fold higher in the Qishon estuary (27 ± 14 × 10^8 cells L^-1) than at the coastal station (Figure 3D). Additionally, BA was found to gradually increase with distance from the coastal environment ($R^2 = 0.75$, $P = 0.008$, $n = 7$), peaking at the bottom of station E3 (Figure 3D and Figure S1C). Often BA comprises a mixture of bacteria with high nucleic acid (HNA) content and bacteria with low nucleic acid (LNA) content (Gasol and del Giorgio, 2000). HNA cells are usually larger and more active than LNA cells (Gasol and del Giorgio, 2000; Servais et al., 2003). In the Qishon estuary and the coastal station, HNA bacteria cells were 2–3-fold more abundant than LNA cells. These high ratios between HNA to LNA cells (>1) provide indirect indication that most of the bacterial community was highly active during the sampling period.

Along the Qishon estuary, bacterial production rates (BP) ranged from 74 to 121 μg C L^-1 d^-1, and were significantly higher (4–7-fold) than at the coastal station (Figure 3E). Similar to phytoplankton and BA, BP were also positively correlated with distance from the coastal environment ($R^2 = 0.83$, $P = 0.003$, $n = 7$). All the above indicate that the microbial community was highly active throughout the estuary (Figure S1D). In fact, BP rates measured in the Qishon estuary were greater than most estuary systems (Ducklow and Carlson, 1992; Meon and Amon, 2004; Apple et al., 2006; Barrera-Alba et al., 2009; Santos et al., 2014), but were in agreement with rates reported from Pensacola Bay estuary in northwestern Florida (Murrell, 2003) and the Hudson River estuary (Findlay et al., 1991). Surprisingly, bacterial production per cell remained constant throughout the different sampling stations, ranging between 28 and 44 fg C cell^-1 d^-1. The sharp rise in BP and BA through the Qishon estuary, while retaining constant BP per cell, implies that the carbon assimilation rate reached a maximum of 36 ± 7 fg C cell^-1 d^-1 for the average bacterial cell.

β-gluconidase (β-glu) is one of various enzymes that are secreted by bacteria (hence defined as an ectoenzyme) to hydrolyze polysaccharides such as TEP into smaller, bio-available macromolecules (Radić et al., 2006; Engel et al., 2014). Similar to TEP, β-glu hydrolysis rates were significantly higher (2–3-fold) in all estuary samples than the coastal reference station and were positively correlated ($R^2 = 0.88$, $P = 0.002$, $n = 7$) with distance from the coast (Figure 3F and Figure S1E). For all estuary sampling points, β-glu comprised 30% of the total assimilated carbon measured as BP, indicating on the importance of polysaccharide hydrolysis as a new-bioavailable carbon source in this system. Additionally, significant positive correlations were observed between TEP to bacterial abundance ($R^2 = 0.95$, $P < 0.001$, $n = 7$) and β-glu activity ($R^2 = 0.93$, $P < 0.001$, $n = 7$) (Figures S2A, B). We elucidate that this newly formed bioavailable carbon may than assimilated by bacteria according to cellular requirements. Further, the high hydrolysis rates by β-glu and the significant contribution to assimilated carbon (BP) indicate the importance of polysaccharides to bacterial metabolism in the Qishon estuary.

From the field observations, we conclude that high BA and TEP concentrations in the Qishon estuary increased the probability of polysaccharide-hydrogels to be colonized by bacteria and form bio-aggregates. The significant contribution of β-gluc and the positive correlations with TEP are all indirect indications that these polysaccharides were hydrolyzed to form an additional carbon source for heterotrophic bacterial proliferation.

**TEP Contribution to Bacterial Activity in Confined Bottle Incubations**

The direct link between TEP and heterotrophic bacterial metabolism was further characterized by carrying out 18 bottle-microcosms. To model TEP contribution to bacterial activity, gum xanthan (GX) was added to nine “enriched” bioassays as pure polysaccharide substrate, since the chemical composition of naturally formed TEP varies drastically (Passow, 2002). Changes in BA, heterotrophic bacterial composition (LNA and HNA), metabolic activity, TEP concentrations and ectoenzymatic hydrolysis rates were measured at the beginning and end of the incubation. Values from the control (un-amended estuary water) incubations were than compared to the enriched (GX) microcosms.

Water samples (1L) were collected from the surface and the bottom of stations E1–E3 along the Qishon estuary, during the summer months (July and August, 2014). To focus on the bacterial community, sampled water were pre-filtered (1 μm) to remove any large suspended particles and micro-phytoplankton. As a result, most phytoplankton were dismissed, evident by the drastic reduction (~90%) in Chl a concentrations, while most heterotrophic bacterioplankton were retained (70–90%). Additionally, it is likely that most grazers such as copepods (e.g., *Acartia* sp.) and isopods (e.g., *Cirolana* sp.) were also removed by the pre-filtration step. To prioritize bacterial heterotrophs while maintaining the native physicochemical conditions of our study site, all microcosm bottles were incubated under dark conditions for 2 days ($T_{2d}$).

**Un-amended, Control Bioassays**

Over the course of the incubations ($T_0$–$T_{2d}$), phytoplankton decreased by 83 ± 17%, while BA and corresponding heterotrophic bacterial biomass (HBB) increased by 32 ± 31% (Figure 4A, Table 2). During this time, the heterotrophic bacterial community showed no change in the HNA/LNA ratio (~2) compared to the initial ($T_0$) conditions (Figure 5, Table 2). Although BA was high at the end of the control incubations in respect to initial concentrations, growth rates were exceptionally low (Figure 4B, Table 2) compared to other marine and fresh water environments (Hamasaki et al., 2004; Smith and Benner, 2005; Azam and Malfatti, 2007; Barrera-Alba et al., 2009).

Concurrently, initial TEP concentrations increased by up to 2.4 ± 1-fold resulting in polysaccharide accumulation, ΔTEP; $T_{2d}$-$T_0 > 0$ (Figure 4C, Table 2). Further, TEP concentrations were found to positively correlate with the increase in heterotrophic bacterial biomass (Figure 6). The accumulation of TEP, the positive correlation with heterotrophic bacterial growth (ΔHBB; $T_{2d}$-$T_0 > 0$) and the extremely low phytoplankton...
FIGURE 4 | Differences between the control and GX-enriched microcosm experiments comparing: changes in heterotrophic bacterial biomass \( \Delta HBB = T_2 - T_0 \) (A), growth rates (B) and TEP concentrations \( \Delta TEP = T_2 - T_0 \) (C) as well as \( \beta \)-glucosidase activity \( \beta \)-glu (D), bacterial carbon demand (BCD) (E), and bacterial growth efficiency (BGE) (F). Each box plot represents microcosms that were incubated with water samples from the surface and bottom of stations E1–E3. Box plot boundary are the 25th–75th percentile while the inner line represent median and whiskers highlight the 10th–90th percentile. Significant differences between control and GX enriched microcosms are marked with an asterisk \( P < 0.01 \).

concentrations, all indicate that in the control incubations TEP was mostly generated by the heterotrophic bacterial community. We suggest that some of the polysaccharide secretions were triggered by viruses that passed filtration and may infect the bacterial community as previously suggested for phytoplankton (Vardi et al., 2012; Lønborg et al., 2013). Concurrent to these direct bacterial TEP secretions, TEP precursors (5–400 nm) that passed the filtration step may self-assembled during the incubation by diffusion and electrostatic interaction to form larger, new TEP (Chin et al., 1998; Verdugo, 2012; Bar-Zeev et al., 2015).

Polysaccharide hydrolysis by \( \beta \)-glucosidase activity \( \beta \)-glu ranged between 20 and 47 \( \mu \)g C L\(^{-1}\) d\(^{-1}\) (Figure 4D), using a conversion factor of 72 \( \mu \)g C to 1 \( \mu \)M MUF (Hoppe, 1993). These ectoenzyme activity rates were relatively high compared to those observed in other eutrophic and oligotrophic environments (Rath et al., 1993; Martínez et al., 1996; Radić et al., 2006). However, they were lower than reported around oil slicks as high hydrocarbon source (Ziervogel et al., 2014). Cell specific hydrolysis rates \( (12 \pm 5 \text{ fg C cell}^{-1} \text{ d}^{-1}) \) were also higher than values reported from the open ocean (Baltar et al., 2009), but highly comparable to hydrolysis rates by bacteria that harbored aggregates (Karner and Herndl, 1992; Azúa et al., 2003; Corinaldesi et al., 2003).

Bacterial respiration (BR) and bacterial production (BP) have varied between the nine incubations, ranging from 45 to 489 \( \mu \)g C L\(^{-1}\) d\(^{-1}\) and 86 to 162 \( \mu \)g C L\(^{-1}\) d\(^{-1}\), respectively. Carbon oxidation through BR (measured as oxygen reduction) was determined following two key assumptions: (i) since most phytoplankton were removed through size exclusion and hindered by the darken incubations, we estimated that community respiration was ascribed mostly (\( \sim 90\% \)) to microbial heterotrophs. (ii) Converting oxygen consumption to carbon utilization was done by a conservative respiratory-quotient (RQ) of 1 (del Giorgio and Cole, 1998; Anesio et al., 2003; Smith and Prairie, 2004).

Heterotrophic metabolism rates were derived from both anabolic (BP) and catabolic (BR) reactions, hereafter defined as bacterial carbon demand \( (\text{BCD} = \text{BP} + \text{BR}) \) and bacterial growth efficiency \( (\text{BGE} = \text{BP}/\text{BCD}) \). In the control incubations, calculated BCD was 359 \( \pm 167 \) \( \mu \)g C L\(^{-1}\) d\(^{-1}\) and BGE values were 42 \( \pm 22\% \), \( n = 9 \) (Figures 4E,F). BCD corresponds to...
TABLE 2 | Biological properties measured in un-amended controls and gum-xanthan (GX) enriched microcosms incubated 2 days (T_2d) under dark + DCMU conditions.

| Parameter | Units | Control | GX enriched |
|-----------|-------|---------|-------------|
|           |       | T_0     | T_2d        | T_0          | T_2d         |
| Chl. a    | µg L⁻¹ | 2.1 ± 0.6 | 0.3 ± 0.2 | 2.1 ± 0.6 | 0.3 ± 0.3 |
| HBB⁸        | µg C L⁻¹ | 47 ± 22 | 60 ± 29 | 47 ± 22 | 86 ± 44 |
| HNA         | x10⁸ Cell L⁻¹ | 15 ± 7.1 | 20 ± 10 | 15 ± 7.1 | 31 ± 14 |
| LNA         | x10⁸ Cell L⁻¹ | 8.3 ± 4.6 | 10 ± 4.2 | 8.3 ± 4.6 | 11 ± 6.3 |
| HNA/LNA     | Cell/Cell | 2 ± 0.7 | 1.9 ± 0.6 | 2 ± 0.7 | 3.3 ± 1.3 |
| TEP         | µg C L⁻¹ | 160 ± 48 | 384 ± 206 | 1312 ± 52 | 849 ± 351 |
| β-glu⁵      | µg C L⁻¹ d⁻¹ | 34 ± 7.5 | 32 ± 9 | 34 ± 7.5 | 46 ± 9 |
| BP¹         | µg C L⁻¹ d⁻¹ | 109 ± 21 | 165 ± 34 | 109 ± 21 | 121 ± 24 |
| BCD         | µg C L⁻¹ d⁻¹ | NA | 359 ± 167 | NA | 557 ± 276 |
| BGE         | % | NA | 42 ± 22 | NA | 37 ± 20 |
| Growth rate⁶ | d⁻¹ | 0.15 ± 0.1 | NA | 0.28 ± 0.09 |

All values are the averages and their standard deviation of nine bottle incubations.

⁸ Bacterial abundance was converted into biomass (HBB) by a factor of 20 fg C per cell (Lee and Fuhrman, 1987).
⁵ TEP was converted from GX into carbon biomass using a 0.74 factor (Engel and Passow, 2001).
⁵ β-glucosidase activity (β-glu) was converted from MUF to carbon biomass using 72 µg C to 1 µM MUF factor (Hoppe, 1993).
⁶ Bacterial production was converted into carbon biomass using a conversion factor of 3.1 kg C mol⁻¹ with an isotope dilution factor of 2.0 to calculate BP (Simon and Azam, 1989).
⁶ Growth rates calculated according to Equation (1).

FIGURE 5 | Changes in heterotrophic bacterial community composition following GX addition (GX) compared to un-amended controls (C) during the experimental microcosm 2 days incubations (T₀–T₂). The bacterial community was segregated into two subgroups containing high and low nucleic acid content (HNA and LNA, respectively). Each box plot represents microcosms that were carried with water samples from the surface and bottom of stations E1–E3. Differences between HNA and LNA were evaluated using One-Way ANOVA followed by Fisher LSD multiple comparison post-hoc test with a confidence of 95%.

FIGURE 6 | Relationship between changes in TEP concentrations (ΔTEP = T₂-T₀) and differences in heterotrophic bacterial biomass (ΔHBB = T₂-T₀). Data was collected from both the control (white circles) and GX enriched (gray circles) bottle incubations. Linear regression is shown of: ΔTEP vs. ΔHBB for the control (R² = 0.54, P = 0.014, n = 9) and enriched (R² = 0.40, P = 0.09, n = 9) microcosms.

the carbon biomass that is required to sustain the net bacterial metabolic needs, while BGE values point on the efficiency at which carbon is assimilated into bacterial biomass (del Giorgio and Cole, 1998; Ducklow et al., 2000). Since BP was determined by short (4 h) incubations, while BR was estimated by measuring oxygen consumption over 48 h, BCD and BGE may have been overestimated to some extent. Further bias might result from variations in bacterial community composition (e.g., HNA: LNA) and by using a conservative RQ of 1, whereas values are known to vary between 0.5 and 4, depending on the oxidation potential of organic compounds (Berggren et al., 2012). Nevertheless, both BCD and BGE measured in this study (neglecting two, high outgroups) were highly comparable with other eutrophic, marine and fresh water environments (Apple and del Giorgio, 2007; Bhaskar and Bhosle, 2008; Amado et al., 2013; Ziervogel et al., 2014).

We suggest that although heterotrophic bacterial abundance (BA) was higher at the end of the control incubations than initial values, the low growth rates relative to other
environments indicate that metabolic activity was mostly
tunnelled to maintain cellular energetic requirements rather
than growth and propagation. These results imply that even
in a eutrophic environment such as the Qishon estuary,
bacterial heterotrophs were possibly nutrient (e.g., carbon)
deficient.

**Gum Xanthan (GX) Enriched Bioassays**

Following GX additions (comprising glucose, mannose and
glucuronic acid), TEP concentrations increased by 8-fold
compared to the control (un-amended) microcosms (Table 2).
By the end of the incubations (T0–T2d), heterotrophic bacterial
growth rates doubled compared to the control, resulting in
80 ± 40% increase in BA (Figures 4A,B, Table 2). The raise
in BA was mostly attributed to drastic HNA proliferation
(Figure 5, Table 2). HNA bacterial activity was shown in both
laboratory experiments (Servais et al., 2003) and elevated
hydrolysis of carbohydrates in field studies (Piontek et al., 2014)
highlighting the potential of HNA to proliferate in carbon
enriched systems.

Compared to the control incubations, ∆HBB (T2d- T0)
significantly increased following GX enrichment, while TEP
was reduced (∆TEP < 0), pointing on intense polysaccharide
hydrolysis (Figures 4A,C). Moreover, the negative correlation
found between ∆TEP and ∆HBB (Figure 6) is an additional
indication that heterotrophic bacterial growth was linked to TEP
consumption.

We suggest that following the addition of GX, TEP
centrations decreased through enhanced ectoenzymatic
(β-gluc) activity relative to the initial rates, T2 > T0; 36 ± 17% (Table 2). Further, at the end of the enriched incubations, β-gluc activity was significantly higher than measured at the control bottles (Figure 4D). These intense hydrolysis rates by β-gluc were likely triggered by the polysaccharide (GX) substrates, as well as the close proximity of the ectoenzymes released by bacteria harboring TEP (Simon et al., 2002; Azam and Malfatti, 2007;
Stocker, 2012). We determined that this newly formed carbon
comprised 51 ± 43% of the heterotrophic BCD in the nine
enriched bioassays. This bioavailable source of carbon could
then prompt bacterial growth by enhancing both anabolic and
catabolic reactions (Figure 4E). High BCD, through positive
feedback-loops may result in constant polysaccharide release,
forming TEP that could then be hydrolyzed as fresh, available
carbon source.

In contrast to BCD, no changes in average BGE (using RQ of 1)
were measured following GX addition compared to the control
incubations (Figure 4F), since both BP and BR increased in a
similar ratio (40 and 50% respectively). We do acknowledge that
variations in the organic composition may shift RQ and alter
BGE. However, adjusting RQ to 1.2 as previously suggested for
glucose (Berggren et al., 2012) did not significantly change BGE
values.

From all the confined incubations we deduced that GX
addition as newly formed polysaccharide-rich substrates were
most likely harbored by bacteria forming “hotspots” with
enhanced BCD. We suggest that the high metabolic requirements
were paired with intense ectoenzymatic (e.g., β-gluc) activity,
hydrolyzing polysaccharide chains into bioavailable organic
compounds (Hoppe, 1993; Martinez et al., 1996; Simon
et al., 2002; Piontek et al., 2014). The consequence of that
newly introduced carbon source, even in a highly eutrophic
environment such as the Qishon estuary was heterotrophic
microbial proliferation. Yet, it is possible that once carbon
is available, other nutrient might become limiting, thereby
specialized heterotrophic bacteria such as diazotrophs could
utilize these new conditions (e.g., high C > N) and prevail (Rahav
et al., 2013).

**Conclusion**

Our study, carried out at the field and through bottle-bioassays,
was designed to link TEP to heterotrophic metabolism and
bacterial growth in a eutrophic estuary system. Along the Qishon
estuary TEP concentrations gradually increased, concurrently
with the development of phytoplankton and bacterioplankton
biomass, most probably through active excretion of polymeric
substances (Figure 3). TEP were often found as large bio-
aggregates, scaffolding algae, bacteria and detritus matter
(Figure 3C). The tight association between TEP, as organic
substrate, and bacteria has facilitated polysaccharide hydrolysis
through intense β-gluc-ectoenzymatic activity (Figures 3, 4).

We postulate that this newly bioavailable carbon source is
formed and consumed along the Qishon estuary through
a positive feedback loop: (i) initially, bacterial anabolic and
catabolic reactions are prompt, thereby increasing heterotrophic
growth rates, mainly of the HNA subgroup (Figure 5). (ii)
While proliferating, bacteria constantly secrete dissolved and
particulate organic matter that forms new TEP, which may later
be hydrolyzed.

Based upon the results above we deduce that heterotrophic
bacteria were limited by carbon, despite the eutrophic nature
of the Qishon estuary. Hence, regardless of the carbon source
(autotrophic, heterotrophic or anthropogenic), when fresh TEP
(e.g., GX) are introduced to the Qishon eutrophic-water, the
bacterial community proliferates. Currently, we cannot rule-out
co-limitation by other microelements such as phosphorous
and/or nitrogen once carbon is supplemented. Regardless, it is
clear that in the Qishon estuary and likely in similar systems,
TEP act as a dynamic metabolic link fueling the microbial
loop, concurrently to other possible roles such as aggregation,
sedimentation and pathogen transfer.

**Author Contributions**

EB and ER designed the research; ER conducted the experiments;
EB and ER analyzed the data; EB and ER wrote the paper.

**Acknowledgments**

This work was partly supported by the PERSEUS project (EC
Contract #287600 to BH). This research was made possible by
the postdoctoral fellowship (to EB) provided by the United
States-Israel Binational Agricultural Research and Development
Supplementary Material

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2015.00403/abstract
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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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