Structural Studies of the Lipopolysaccharide from the Fish Pathogen Aeromonas veronii Strain Bs19, Serotype O16

Anna Turska-Szewczuk 1,* , Katarzyna A. Duda 2, Dominik Schwudke 3, Agnieszka Pekala 4, Alicja Kozinska 4 and Otto Holst 2

1 Department of Genetics and Microbiology, M. Curie-Sklodowska University, Akademicka 19, Lublin 20-033, Poland
2 Division of Structural Biochemistry, Research Center Borstel, Leibniz-Center for Medicine and Biosciences, Parkallee 4a/c, Borstel D-23845, Germany; E-Mails: kduda@fz-borstel.de (K.A.D); oholst@fz-borstel.de (O.H.)
3 Division of Bioanalytical Chemistry, Research Center Borstel, Leibniz-Center for Medicine and Biosciences, Parkallee 10, Borstel D-23845, Germany; E-Mail: dschwudke@fz-borstel.de
4 Department of Fish Diseases, National Veterinary Research Institute, Partyzantow 57, Pulawy 24-100, Poland; E-Mails: a.pekala@piwet.pulawy.pl (A.P.); koala@piwet.pulawy.pl (A.K.)

* Author to whom correspondence should be addressed; E-Mail: aturska@hektor.umcs.lublin.pl; Tel.: +48-81-537-50-18; Fax: +48-81-537-59-59.

Received: 24 December 2013; in revised form: 27 January 2014 / Accepted: 8 February 2014 / Published: 7 March 2014

Abstract: Chemical analyses, mass spectrometry, and NMR spectroscopy were applied to study the structure of the lipopolysaccharide (LPS) isolated from Aeromonas veronii strain Bs19, serotype O16. ESI-MS revealed that the most abundant LPS glycoforms have tetra-acylated or hexa-acylated lipid A species, consisting of a bisphosphorylated GlcN disaccharide with an AraN residue as a non-stoichiometric substituent, and a core oligosaccharide composed of HepsHex3HexN2Kdo1P1. Sugar and methylation analysis together with 1D and 2D 1H and 13C NMR spectroscopy were the main methods used, and revealed that the O-specific polysaccharide (OPS) of A. veronii Bs19 was built up of tetrasaccharide repeating units with the structure: →4)-α-D-Quip3NAc-(1→3)-α-L-Rhap-(1→4)-β-D-Galp-(1→3)-α-D-GalpNAc-(1→. This composition was confirmed by mass spectrometry. The charge-deconvoluted ESI FT-ICR MS recorded for the LPS preparations identified mass peaks of SR- and R-form LPS species, that differed by Δm = 698.27 u, a value corresponding to the calculated molecular mass of one OPS repeating unit (6dHexNAc6dHexHexHexNAc-H2O). Moreover, unspecific fragmentation spectra confirmed
the sequence of the sugar residues in the OPS and allowed to assume that the elucidated structure also represented the biological repeating unit.

**Keywords:** lipopolysaccharide; $O$-specific polysaccharide; *Aeromonas veronii*; fish pathogen; ESI MS; NMR

**Abbreviations**

NMR, nuclear magnetic resonance; ESI-MS, electrospray ionization mass spectrometry; FT-ICR, Fourier transform ion cyclotron resonance; OPS, $O$-specific polysaccharide; ADP, adenosine diphosphate; SDS-PAGE, sodium dodecyl sulfate polyacrylamide gel electrophoresis; ESI-FT-ICR, electrospray ionization Fourier transform ion cyclotron resonance; GC-MS, gas chromatography with mass spectrometry; DQF-COSY, double quantum filtered correlation spectroscopy; NOE, Nuclear Overhauser effect; ROESY, rotating frame Overhauser effect spectroscopy; TOCSY, total correlation spectroscopy; PCR-RFLP, polymerase chain reaction/restriction fragment length polymorphism.

**1. Introduction**

*Aeromonas* spp. bacteria are widespread in aquatic environments and soil habitats and are also frequently isolated from raw and processed food. They are either mesophilic, motile, or psychrophilic non-motile Gram-negative rods [1–3]. *Aeromonas* strains identified as members of the gut microflora in fish and other aquatic animals (amphibians, reptiles) may cause various diseases under environmental stress conditions (overcrowding, poor water quality, organic pollution, and hypoxia) [4,5]. Amongst mesophilic and motile species, *A. hydrophila*, *A. caviae*, *A. sobria*, and *A. veronii* have been described as important fish pathogens. They cause chronic disease with open dermal ulcers and other pathological lesions or acute systemic infection referred to as motile aeromonad septicemia (MAS) [5–7].

These bacteria, especially belonging to the species *A. hydrophila*, *A. caviae*, and *A. veronii* bv. *sobria*, often have been associated with several categories of human infections. Clinical presentations of such diseases comprise both gastrointestinal, frequently foodborne diseases, and life-threatening extraintestinal infections, including septicemia, wound and urinary tract infections, and, occasionally, meningitis, especially in immunocompromised patients and children [5,8–11], however, recent data indicates that *Aeromonas* may also be primary causes of infections in immunocompetent hosts [12].

The possible mechanisms of *Aeromonas* infections are complex and multifactorial. Several extracellular products of *Aeromonas* including hemolysins, cytotonic and cytotoxic enterotoxins, proteases, lipases, and leucocidins have been suggested as possible contributory factors in the pathogenesis of these bacteria [12]. Amongst these, a type II secretion system (secretion of enterotoxin-Act) and a type III secretion system (T3SS) seem to be leading [10]. Recently characterized effectors of the type VI secretion system had actin-ADP ribosylation activity that induced host cell cytotoxicity [5,10].
Moreover, cell-surface components such as outer membrane proteins, lipopolysaccharide (LPS), S-layer, polar flagella, and pili (type IV and bundle-forming pili) have been identified as *Aeromonas* putative virulence factors [13–16]. An equally important non-fimbrial adhesion factors that have been implicated in the pathogenesis of *Aeromonas* spp. are S-layer and LPS. The S-layer enhances certain physical attributes of the bacterium, including increases in cellular hydrophobicity, cell aggregation, and cell-to-tissue adhesion [17]. As an adhesine, S-form LPS is indispensable for initial attachment of bacteria to host tissue and necessary during infection events, where it protects bacteria from complement-mediated killing and antimicrobial peptides [5]. It is plausible that some virulence factors located in the outer membrane require the presence of O-antigens (O-specific polysaccharides, OPS) for proper expression or functionality. In addition, the OPS variations seem to play an essential role at several stages of the infection process, including the adherence step and the ability to protect against host defense mechanisms [5]. Although it was not clearly evidenced which structural determinants are the most important for virulence, it was found that some O serotypes are more frequently associated with certain infections. Studies demonstrated that *Aeromonas* strains belonging to serogroups O11, O16, O18, and O34 (Sakazaki and Shimada scheme [18]) are associated with most cases of bacteremia, implying the OPS variants are relevant in systemic disease pathogenesis [5]. It was proven that smooth LPS was implicated to promote adhesion of *A. veronii* bv. *sobria* to HEp-2 cells, and switching off of the OPS by using specific monoclonal antibodies impaired bacterial adherence [19,20].

The species *A. veronii*, originally described by Hickman-Brenner et al. (1987), as a novel member of the genus is commonly diarrhea-associated and is a rare cause of bacteremia [21]. However, there were also reports of *A. veronii* bv. *veronii* and *A. veronii* bv. *sobria* septicemia incidences in immunocompromised patients [22,23]. Nevertheless, this species is commonly known as fish pathogen, especially associated with ulcerative syndrome [6,7]. In the light of the increased *Aeromonas* infection incidence rate and the economic importance of these diseases in fish farms as well as possible public health effects, it is essential to characterize the virulence factors of these bacteria.

In the majority of the *Aeromonas* strains studied, the OPS has a heteropolymeric repeating unit and contains aminohexoses and amino-6-deoxyhexoses as well as acidic non-carbohydrate substituents such as 3-hydroxy butyrate [24–27]. Recently, two new structures of OPS were established for the species *A. bestiarum*, which strains are frequently isolated in the course of motile aeromonad septicemia in Polish commercial ponds [28–30].

Now we report on the chemical structure of the LPS from *A. veronii* strain Bs19 serologically classified as serogroup O16, which was isolated from skin of carp (*Cyprinus carpio* L.) with hemorrhagic and necrotic ulcers [31].

2. Results and Discussion

2.1. Isolation of LPS and SDS–PAGE

*A. veronii* Bs19 LPS was isolated by hot phenol-water extraction [32] from enzymatically digested bacterial cells [29]. It was found that the LPS species distributed between the water and phenol phases as hydrophilic and hydrophobic fractions, in yields of 4.2% and 0.3% of the dry bacterial cell mass,
respectively. The SDS-PAGE analysis (Figure 1) of these preparations revealed that the smooth, S-form LPS species depicting a typical ladder-like pattern were recovered mainly from the water phase, and the rough R- or semi-rough SR-LPS species from the phenol phase. The yield of the hydrophobic LPS fraction, extracted from the phenol, which was less than 10% of the yield obtained from water, indicated a lower content of R-form LPS molecules in the cell envelope of A. veronii Bs19.

Figure 1. Silver-stained SDS-Tricine PAGE of the water- and phenol-soluble LPS fractions of A. veronii strain Bs19 (lane 2, lane 3, respectively), and Salmonella enterica sv. Typhimurium as reference (lane 1). Two micrograms were loaded per lane.

2.2. Chemical and ESI FT-ICR Mass Spectrometric Analyses of LPS

Sugar analyses of the LPS preparations were performed by GC-MS of the alditol acetates. The R- and SR-form LPS species contained Glc, GlcN, D,D-Hep, and L,D-Hep in a molar ratio of approx. 2.9:0.9:2.6:1. All these sugars (in a molar ratio of approx. 3.3:1:4.0:1) were also found in the fraction obtained from the water phase, which contained high molecular mass S-form LPS species. Additionally, the chemical analysis of both preparations showed 6-deoxyhexose (Rha), 3-amino-3, 6-dideoxyglucose (Qui3N), Gal and GalN, in a molar ratio of approx. 1.2:1:1.2:1.0. These sugars were identified as components of the OPS (see Section 2.3). Kdo (3-deoxy-d-manno-2-octulosonic acid)—the only acidic sugar—was found in both LPS fractions. GC-MS analysis of the fatty acids as methyl esters and O-TMS derivatives identified 3-hydroxy myristic [14:0(3-OH)] and dodecanoic (12:0) acids, as the most abundant species. GlcN was identified as the sugar component of the lipid A.

The LPS preparations from A. veronii Bs19 were analyzed by ESI FT-ICR MS. The charge-deconvoluted ESI MS (negative-ion mode) (Figure 2A,B) of both LPS fractions showed a complex pattern of molecules originating from heterogeneity of lipid A and the core oligosaccharide. The heterogeneity was caused by non-stoichiometric substitutions with hexose (Hex, Δm = 162.05 u), heptose (Hep, Δm = 192.06 u), one or two fatty acid residues, 14:0(3-OH) (Δm = 226.19 u), and 14:0(3-OH) + 12:0 (Δm = 408.36 u), respectively, as well as peaks originating from different acyl chain length (i.e., 12:0 and 14:0).
The mass spectra of LPS preparations showed a molecular peak at 3268.403 u corresponding to a R-LPS glycoform with the core decasaccharide -Hep₂Hex₃HexN₉Kdo₁P₃ linked to tetra-acylated lipid A (LPS₄tetra), which possessed a bisphosphorylated diglucosaminyl backbone (Figure 2, Table 1).

**Figure 2.** Charge-deconvoluted ESI FT-ICR mass spectra (negative ion mode) of the water- and phenol-soluble LPS fractions from *A. veronii* strain Bs19 (A) and (B), respectively. Indicated mass values refer to the monoisotopic signals of the neutral molecules. LPS₄tri, LPS₄tetra, LPS₆hexa, acylation state of the lipid A, *undefined* contaminations.

**Table 1.** Composition of the main species present in the charge deconvoluted ESI FT-ICR mass spectra (negative ion mode) of the water and phenol-soluble lipopolysaccharide (LPS) fractions of *A. veronii* Bs19.

| Species       | M<sub>measured</sub> water phase | M<sub>measured</sub> phenol phase | M<sub>calculated</sub> | Composition                                                                 |
|---------------|---------------------------------|-----------------------------------|------------------------|-----------------------------------------------------------------------------|
| LPS tetra I   | 3042.219                        | 3042.214                          | 3042.203               | Hep₂Hep₂Hep₂Hex₃HexN₉Kdo₃P₃[14:0(3-OH)]₁₂:0                                 |
| LPS tetra II  | 3268.403                        | 3268.400                          | 3268.397               | Hep₂Hep₂Hep₂Hex₃HexN₉Kdo₃P₃[14:0(3-OH)]₁₂:0                                 |
| LPS tetra III | 3676.765                        | 3676.765                          | 3676.757               | Hep₂Hep₂Hep₂Hex₃HexN₉Kdo₃P₃[14:0(3-OH)]₁₂:0                                 |
| SR-LPS tetra I| 3740.498                        | 3740.498                          | 3740.480               | 6dHex₆dHexNHep₂Hep₂Hep₂Hex₃HexN₉Kdo₃P₃Ac₂[14:0(3-OH)]₁₂:0                  |
| SR-LPS tetra II| 3966.695                       | 3966.694                          | 3966.671               | 6dHex₆dHexNHep₂Hep₂Hep₂Hex₃HexN₉Kdo₃P₃Ac₂[14:0(3-OH)]₁₂:0                  |
| LPS hexa I    | 4375.077                        | 4375.077                          | 4375.031               | 6dHex₆dHexNHep₂Hep₂Hep₂Hex₃HexN₉Kdo₃P₃Ac₂[14:0(3-OH)]₁₂:0                  |
| LPS hexa II   | 2816.290                        | 2816.300                          | 2816.292               | Hep₂Hep₂Hep₂Hex₃HexN₉Kdo₃P₃[14:0(3-OH)]₁₂:0-H₂O                           |
| LPS hexa III  | 2624.231                        | 2624.240                          | 2624.231               | Hep₂Hep₂Hep₂Hex₃HexN₉Kdo₃P₃[14:0(3-OH)]₁₂:0-H₂O                           |

Moreover, the spectra showed signals at 3740.498 and 3966.695 u (Figure 2), which corresponded to the semi-rough LPS glycoforms: SR-LPS₄tri I and SR-LPS₄tetra I, respectively, with different acylation patterns of the lipid A and carrying one OPS repeating unit (6dHexNAc₆dHexHexHexNAc) minus
H$_2$O (calculated mass 698.274 u). The measured mass difference of $\Delta m = 698.27$ u was in full accordance with the chemical structure determined by NMR spectroscopy (see Section 2.3). Exclusively, the spectrum of the water-soluble fraction indicated signals at 3676.765 and 4375.077 u attributed to LPS glycoforms with hexa-acylated lipid A, namely $R$-LPS$_{\text{hexaI}}$ and SR-LPS$_{\text{hexaI}}$, respectively. On the other hand, in the mass spectrum of the phenol-soluble LPS, two intensive signals at 2624.231 and 2816.290 u were assigned to the $R$-LPS glycoforms with tetra-acylated lipid A and a shorter core region being hepta- and octasaccharide, respectively (Figure 2, Table 1).

For a more detailed interpretation, both LPS preparations were unspecifically fragmented in the collision cell which yielded Y- and B-mass fragments arising from cleavage of the labile ketosidic linkage between lipid A and Kdo of the core oligosaccharide (Figure 3) [33].

**Figure 3.** Charge-deconvoluted ESI FT-ICR mass spectra (negative ion mode) of the water- and phenol-soluble LPS fractions from *A. veronii* Bs19 with unspecific fragmentation in the collision cell (collision voltage 30 V) (A) and (B), respectively, providing the fragmentation of the labile linkage between the Kdo and lipid A. Mass numbers given refer to the monoisotopic masses of the neutral molecules.

In particular, the mass spectra showed, e.g., signals at 1907.588 and 1809.599 u, which corresponded to the B-fragments originating from the core glycoforms and Y-fragments at 1360.830 and 1134.638 u derived from the lipid A species. In addition to these, a further B-fragment at 2605.869 u was observed, which was attributed the core oligosaccharide with one O-antigen repeating unit attached (Table 2).

Based on the chemical component analysis of main fragments, e.g., Y$^{II}$ and B$^{II}$ as well as Y$^{II}$ and B$^{IV}$, using high-resolution mass spectrometric data, the molecular composition of the LPS glycoforms, $R$-LPS$_{\text{tetraI}}$ and SR-LPS$_{\text{tetraI}}$, respectively, was confirmed.
Table 2. Composition of the main Y- and B-fragments present in the charge deconvoluted ESI FT-ICR mass spectra (negative ion mode) of the water- and phenol-soluble LPS fractions from A. veronii strain Bs19 obtained with unspecified fragmentation.

| Species | M_{measured}^{water phase} | M_{measured}^{phenol phase} | M_{calculated} | Composition |
|---------|---------------------------|-------------------------------|----------------|-------------|
| Y¹      | 1134.638                  | 1134.639                     | 1134.634       | HexN₃P₂[14:0(3-OH)]₁₂:0  |
| Y₁      | 1162.673                  | 1162.672                     | 1162.665       | HexN₃P₂[14:0(3-OH)]₁₄:0  |
| Y²      | 1360.830                  | 1360.830                     | 1360.827       | HexN₃P₂[14:0(3-OH)]₁₂:0  |
| Y²      | 1388.831                  | 1388.832                     | 1388.859       | HexN₃P₂[14:0(3-OH)]₁₄:0  |
| Y²      | 1769.193                  | –                             | 1769.188       | HexN₃P₂[14:0(3-OH)]₁₂:0  |
| B¹      | 1809.599                  | 1809.598                     | 1809.598       | Hep₃Hex₃HexNKdo-2H₂O      |
| B²      | 1907.588                  | 1907.588                     | 1907.569       | Hep₃Hex₃HexNKdo-P₃H₂O     |
| B²      | 2507.880                  | 2507.875                     | 2507.866       | 6dHex6dHexNHex₃HexN_KdoAc₂-2H₂O |
| B²      | 2605.869                  | 2605.876                     | 2605.843       | 6dHex6dHexNHex₃HexN_KdoP₂Ac₂-2H₂O |

2.3. Structural Studies of the OPS

The OPS was released from the water-soluble LPS fraction by mild-acid degradation followed by gel-permeation chromatography (GPC). Sugar analysis of the OPS (GC-MS of the alditol acetates) revealed Rha, Qui₃N, Gal, and GalN in a relative peak area ratio of approx. 1:0.9:1.1:1.2. Qui₃N was identified by comparing its retention time and mass spectra with those compounds obtained from the OPS of E. coli O5 [34] (a strain kindly provided from the Institute of Immunology and Experimental Therapy, Wroclaw, Poland) and Aeromonas veronii bv. sobria strain K49 [35]. The absolute configuration of monosaccharides determined by GLC of the acetylated (S)-2-butyl glycosides [36] identified the configuration of Rha as L and that of the other sugars as D.

Linkage analysis by GC-MS of the partially methylated alditol acetates derived from the methylated polysaccharide resulted in identification of 3-substituted Rhap, 4-substituted Qui₃N, 3-substituted GalpN, and 4-substituted Galp.

The ¹H and ¹³C NMR spectra showed that the OPS of A. veronii strain Bs19 had a regular structure composed of linear tetrasaccharide repeating units. The ¹H NMR spectrum of the OPS (Figure 4A) contained signals for four anomeric protons at δ 5.27, 5.21, 5.05, and 4.47, labeled A through D, respectively. In the high field region of the spectrum, there were also signals originating from the methyl groups of 6-deoxysugars (Rha and Qui₃N) at δ 1.33 and 1.30, and two signals of N-acetyl groups at δ 2.02 and 1.97.

The ¹³C NMR spectrum (Figure 4B) of the OPS contained signals for four anomeric carbons at δ 96.03 (C), 98.11 (A), 102.46 (B), and 106.02 (D); signals for two nitrogen-bearing carbons at δ 49.15 and 55.09 (GalN C-2 and Qui₃N C-3, respectively); two methyl groups of 6-deoxysugars at δ 18.0 and 19.0 (Rha and Qui₃N); N-acetyl groups (CH₃ at δ 23.7 and CO at δ 175.6, and δ 23.2 and CO at δ 175.4, for GalN and Qui₃N, respectively), and other non-anomeric sugar ring carbons in the region δ 68.24–78.33, some of which overlapped. The ¹³C NMR data showed that all the sugar residues were in the pyranose form [37] as no signals for ring carbons above δ 80, diagnostic of furanose, were detected.
Figure 4. $^1$H NMR (700.43 MHz) (A) and $^{13}$C NMR (176.14 MHz) (B) spectra of the OPS from A. veronii strain Bs19. Capital letters and Arabic numerals refer to atoms in the sugar residues denoted as shown in Table 3. NAc, N-acetyl groups, spectra were recorded at 32 °C in D$_2$O as a solvent, relative to external acetone as reference ($\delta$H 2.225, $\delta$C 31.45).

The anomeric configuration of each monosaccharide was assigned on the basis of the $^3J_{H-1,H-2}$ (measured on the DQF-COSY spectrum) and $^1J_{C-1,H-1}$ coupling constants, and the intra-residual NOE contacts identified in the ROESY spectrum, whereas the ring configuration of each residue was inferred by the vicinal $^3J_{H,H}$ coupling constants [37].

Chemical shifts of each spin system were assigned in $^1$H,$^1$H, TOCSY, DQF-COSY, ROESY, $^1$H,$^{13}$C HSQC, and $^1$H,$^{13}$C HMBC experiments. All chemical shifts are summarized in Table 3.

Based on these data, the spin systems were assigned to four residues, one Rha$p$, one Quip3NAc, one Galp, and one GalpNAc. In particular, the spin systems A ($^3J_{1,2}$ ~3.2 Hz) and D ($^3J_{1,2}$ ~8 Hz) were identified as $\alpha$-GalpNAc and $\beta$-Galp residues, respectively [38]. A small $J_{1,2}$ coupling constant ~3.5 Hz and chemical shifts of H-1 ($\delta$ 5.05) and C-5 ($\delta$ 68.24) indicated that the spin system C was $\alpha$-linked Quip3NAc [39]. The chemical shifts for H-5 and C-5 at $\delta$H 3.78 and $\delta$C 70.71, respectively, as well as signals of H-6 at $\delta$H 1.33 from the methyl group indicated that the spin system B was $\alpha$-linked Rhap [40,41].
Table 3. $^1$H and $^{13}$C NMR chemical shifts of the constituents of the O-specific polysaccharides (OPS) of A. veronii strain Bs19. Spectra were recorded in D$_2$O relative to external acetone as reference ($\delta_H$ 2.225, $\delta_C$ 31.45).

| Sugar Residue | H-1 (C-1) | H-2 (C-2) | H-3 (C-3) | H-4 (C-4) | H-5 (C-5) | H-6 (C-6) |
|---------------|-----------|-----------|-----------|-----------|-----------|-----------|
| $\rightarrow$3)-$\alpha$-D-GalpNAc-(1$\rightarrow$A | 5.27 | 4.35 | 3.95 | 4.24 | 4.11 | 3.75 |
| | 98.11 | 49.15 | 73.32 | 69.72 | 72.30 | 62.30 |
| $\rightarrow$3)-$\alpha$-L-Rhap-(1$\rightarrow$B | 5.21 | 4.28 | 3.87 | 3.61 | 3.78 | 1.33 |
| | 102.46 | 68.34 | 77.14 | 71.21 | 70.71 | 18.00 |
| $\rightarrow$4)-$\alpha$-D-Quip3NAc-(1$\rightarrow$C | 5.05 | 3.56 | 4.33 | 3.52 | 4.20 | 1.30 |
| | 96.03 | 71.50 | 55.09 | 77.44 | 68.24 | 19.00 |
| $\rightarrow$4)-$\beta$-D-Galp-(1$\rightarrow$D | 4.47 | 3.54 | 3.77 | 4.03 | 3.74 | 3.75 |
| | 106.02 | 71.45 | 74.17 | 77.04 | 76.06 | 62.30 |

Chemical shifts for NAc were $\delta_H$ 2.02 and $\delta_C$ 23.7/175.6 for A and $\delta_H$ 1.97 and $\delta_C$ 23.2/175.4 for C.

The galacto configuration of A and D was determined by the small $^3J_{3,4}$ ($\sim$3 Hz) and $^3J_{4,5}$ ($\sim$1 Hz) coupling constants [42,43]. In the TOCSY spectrum, correlations were visible between H-1 and H-2, H-3, H-4, and the other proton signals were assigned by connectivities identified in the ROESY (strong H-3/H-5) and COSY spectra. The $\alpha$-configuration of GalpN A was proven by the intra-residue H-1/H-2 connectivity observed in the ROESY spectrum. In addition, a N-acetamido sugar was confirmed by correlation of H-2 at $\delta$ 4.35 to the corresponding carbon-bearing nitrogen at $\delta$ 49.15, as revealed by the HSQC experiment. The $\beta$-anomeric configuration of Galp D was inferred from the $^1J_{C,H}$ coupling constant (162 Hz) [43] and confirmed by the intra-residue H-1/H-3 and H-1/H-5 correlations observed in the ROESY spectrum.

The ring gluco configuration of the spin system C was assigned according to the large $J_{2,3}$, $J_{3,4}$, and $J_{4,5}$ coupling constant values ($\sim$10 Hz), and was confirmed by correlations of each H-1, to H-6 with all other protons of the residue in the TOCSY spectrum [39]. Moreover, the $\alpha$-anomeric configuration of Qui3N C was also inferred from the $^1J_{C,H}$ coupling constants (173–174 Hz), and from the intra-residue H-1/H-2 correlation observed in the ROESY spectrum. The site of attachment of the amide-bond acetyl group to Qui3N C was confirmed by the correlation of its C-1 with H-3 of the carbon-bearing nitrogen at $\delta_C/\delta_H$ 175.4/4.33.

The manno configuration of B was indicated by relatively high coupling constant values of $^3J_{3,4}$ and $^3J_{4,5}$ ($\sim$10 Hz) contrasting with the small value of $^3J_{2,3}$ $\sim$3.5 Hz [39,44]. For the manno spin system cross-peaks between H-1 and H-2 and H-2 and H-3/H-6, as well as between H-6 and H-1/H-4, were observed in the TOCSY spectrum. The $\alpha$-configuration of Rhap B was also proven by the intra-residue H-1/H-2 connectivity observed in the ROESY spectrum [41].

Low-field positions of the signals for C-3 of $\alpha$-GalpNAC A ($\delta$ 78.33), C-3 of $\alpha$-Rhap B ($\delta$ 77.14), C-4 of $\alpha$-Qui3NAC C at $\delta$ 77.44 and C-4 of $\beta$-Galp D at $\delta$ 77.04, as compared with the chemical shifts of the corresponding non-substituted monosaccharides, elucidated the glycosylation pattern of the sugar residues [37,45].

The sequence of the sugar residues in the repeating unit was determined by $^1$H, $^1$H ROESY and $^1$H,$^{13}$C HMBC experiments. In the 2D ROESY spectrum of the OPS, the following strong NOE
contacts were observed: α-GalpNAc H-1 (A), α-Quip3NAc H-4 (C) at δ 5.27/3.52; α-Quip3NAc H-1 (C), α-Rhap H-3 (B) at δ 5.05/3.87; α-Rhap H-1 (B), β-Galp H-4 (D) at δ 5.21/4.03; and β-Galp H-1 (D), α-GalpNAc H-3 (A) at δ 4.47/3.95. These data were confirmed by a 2D H-detected heteronuclear multiple-bond (1H, 13C HMBC) experiment (Figure 5), which showed the following inter-residue cross-peaks: α-GalpNAc H-1 (A), α-Quip3NAc C-4 (C) at δ 5.27/77.44; α-Quip3NAc H-1 (C), α-Rhap C-3 (B) at δ 5.05/77.14; α-Rhap H-1 (B), β-Galp C-4 (D) at δ 5.21/77.04; and β-Galp H-1 (D), α-GalpNAc C-3 (A) at δ 4.47/78.33.

The data presented above were consistent with mass spectrometry analysis of the lower molecular mass fraction of degraded polysaccharide (PS) isolated from the water-soluble LPS after mild acid hydrolysis and separation using GPC.

**Figure 5.** Section of the 1H,13C HMBC spectrum of the OPS of A. veronii strain Bs19. Correlations between anomeric protons and carbons at the glycosidic linkages are underlined. Some other H/C correlations are also depicted (italic). Capital letters and Arabic numerals refer to atoms in the sugar residues denoted as shown in Table 3.

The charge-deconvoluted ESI FT-ICR mass spectrum obtained with unspecific fragmentation (collision voltage 5 V) resulted in the cleavage of glycosidic linkages and yielded Z and C series of mass fragments, which contained the reducing and terminal end of the molecule, respectively. The Z mass fragments, seen as the pairs of mass peaks that differed by Δm = 18 u (loss of water), were attributed to the Kdo containing part of the molecule. As already described, this was expected due to the formation of anhydro-Kdo forms during mild acid hydrolysis [46]. In particular, Z10, Z14, and Z18 mass fragments at 1827.612, 2525.895 and 3224.159 u corresponded to the calculated molecular mass of the core oligosaccharide (Hep5Hex3HexNKdo-H2O) without or with one and two OPS repeats attached, respectively. In turn, the C series of fragments were ascribed to the O-repeats containing part of the molecule. In detail, the C4 mass fragment at 716.286 u referred to the molecular mass of one OPS repeat (6dHexNAc6dHexHexHexNAc), in turn the C5-C12 fragments allowed to follow up the
sequence of the sugar residues. In summary, the C and Z series of mass peaks confirmed the structure of the OPS established by NMR and enabled to propose the composition of its biological repeating unit. The MS results are shown below (Figure 6, Table 4).

**Figure 6.** (A) Part of the charge-deconvoluted ESI FT-ICR mass spectrum (negative ion mode) of the lower molecular mass fraction of the degraded PS isolated from the LPS of *A. veronii* Bs19, recorded with unspecific fragmentation. (B) Fragmentation scheme of the molecule. Mass numbers given refer to the monoisotopic masses. Mass fragments (marked with capital letters) are labeled according to the nomenclature of Domon and Costello [33].

![Fragmentation Scheme](image)

**Table 4.** Composition of the main species present in the charge deconvoluted ESI FT-ICR MS (negative ion mode) of the lower molecular mass fraction of the degraded PS isolated from the LPS of *A. veronii* Bs19, recorded with unspecific fragmentation. Mass fragments (marked with capital letters) are labeled according to the nomenclature of Domon and Costello [33].

| Species | M$_{measured}$ | M$_{calculated}$ | Composition |
|---------|----------------|-----------------|-------------|
| C$_4$   | 716.286        | 716.284         | [6dHexNAc6dHexHexHexNAc] |
| C$_5$   | 903.372        | 903.369         | [6dHexNAc6dHexHexHexNAc] |
| C$_6$   | 1049.433       | 1049.427        | [6dHexNAc6dHexHexHexNAc]6dHexNAc6dHex |
| C$_7$   | 1211.486       | 1211.479        | [6dHexNAc6dHexHexHexNAc]6dHexNAc6dHexHex |
| C$_8$   | 1414.576       | 1414.558        | [6dHexNAc6dHexHexHexNAc]$_2$ |
| C$_9$   | 1601.669       | 1601.643        | [6dHexNAc6dHexHexHexNAc]$_2$6dHexNAc |
| C$_{10}$| 1747.724       | 1747.700        | [6dHexNAc6dHexHexHexNAc]$_2$6dHexNAc6dHex |
| C$_{12}$| 2112.830       | 2112.831        | [6dHexNAc6dHexHexHexNAc]$_3$ |
| Z$_{10}$| 1827.612       | 1827.603        | Hep$_3$Hex$_2$HexNKdo-H$_2$O |
| Z$_{12}$| 2192.748       | 2192.734        | [HexHexNAc]Hep$_3$Hex$_2$HexNKdo-H$_2$O |
| Z$_{14}$| 2525.895       | 2525.876        | [6dHexNAc6dHexHexHexNAc]Hep$_3$Hex$_2$HexNKdo-H$_2$O |
| Z$_{16}$| 2891.010       | 2891.007        | [6dHexNAc6dHexHexHexNAc]Hep$_3$Hex$_2$HexNKdo-H$_2$O |
| Z$_{17}$| 3037.077       | 3037.065        | [6dHexNAc6dHexHexHexNAc]Hep$_3$Hex$_2$HexNKdo-H$_2$O |
| Z$_{18}$| 3224.159       | 3224.145        | [6dHexNAc6dHexHexHexNAc]Hep$_3$Hex$_2$HexNKdo-H$_2$O |
On the basis of all the data obtained, it was concluded that the biological repeating unit of the OPS from *A. veronii* strain Bs19, serotype O16 had the structure:

\[
\rightarrow 4\)-α-D-Quip3NAc-(1→3)-α-L-Rhap-(1→4)-β-D-Galp-(1→3)-α-d-GalpNAc-(1→
\]

The structure of the OPS from *A. veronii* Bs19 described herein is the first established for the species. A similar tetrasaccharidic backbone, in respect to both sugar sequence and glycosylation pattern, has been identified as the main component of the O-antigen repeating units of *Salmonella Dakar*, serogroup O28, and *Escherichia coli*, serogroup O71 [47,48]. Whereas, in *E. coli* O71, the difference is caused by non-stoichiometric O-acetylation of β-D-Galp, in the OPS of *S. Dakar*, α-d-GalpNAc is additionally substituted at O-4 by β-D-GlcP [47,48]. It is likely that this terminal residue is also responsible for structural differences within the serogroup O28. Some aspects of the molecular basis for subdividing the O28 serogroup of the Kauffmann-White scheme into three subfactors O28, O28₁ and O28₂ has been recently explained [49]. Studies with monoclonal antibodies confirmed that the O28₁—antigen specificity is attributed to the 3-substituted or 3,4-disubstituted α-d-GalpNAc, which constitute the component of the main chain both *S. Dakar* and *Salmonella Telaviv* OPSs [49].

3. Experimental Section

3.1. Bacterial Strain, Cultivation Conditions and Isolation of the LPS

The *A. veronii* strain Bs19 was isolated from pathologically altered skin of carp suffering from ulcerative syndrome in a commercial pond, as previously reported [31], and was obtained from the Collection of the Microorganisms of the Department of Fish Diseases, National Veterinary Research Institute (Pulawy, Poland). Based on both biochemical properties and PCR-RFLP analysis of the 16S rDNA, strain Bs19 was identified to the species level, and classified to the serogroup O16 [28] according to the scheme of Sakazaki and Shimada [18].

The bacteria were cultivated in tryptic soy broth (TSB) at 28 °C for 72 h. The cells were harvested by low speed centrifugation (8500×g, 20 min). The recovered bacterial cell pellet was washed twice with 0.5 M saline and once more with distilled water. Bacterial cells were digested with lysozyme, RNAse, and DNAse (24 h, 1 mg/g), and then with Proteinase K (36 h, 1 mg/g) in 50 mM phosphate buffer (pH 7.0) containing 5 mM MgCl₂. The suspension was dialyzed against distilled water and freeze-dried. The digested cells were extracted three times with aq 45% phenol at 68 °C [32], and the separated layers were dialyzed against deionized water, purified by ultracentrifugation (105,000×g, 4 h) and freeze-dried to give LPS in a yield of 4.5% of dry bacterial cell mass. In total, 890 mg of LPS were isolated, 93.3% of which was *S*-form (water phase) and 6.7% *SR*- and *R*-form (phenol phase). Both LPS preparations were further analyzed.

3.2. Isolation of the OPS

The OPS was obtained by mild acid hydrolysis of the *S*-form LPS (100 mg) with 2% acetic acid at 100 °C for 3 h, followed by GPC of the water soluble-portion on a column (1.8 × 80 cm) of Sephadex G-50 fine (Pharmacia, Sweden) using 1% acetic acid as an eluent and monitoring with a Knauer
differential refractometer (Knauer, Berlin, Germany). The yield of the OPS fraction was 24.5% of the LPS mass subjected to hydrolysis. The sediment released by acid hydrolysis of the LPS was purified by the Bligh-Dyer method as it was described earlier [50] to give 12 mg lipid A.

3.3. Chemical Analyses

For neutral and amino sugar analysis, the LPS samples and the OPS were hydrolyzed with 2 M CF$_3$CO$_2$H (120 °C, 2 h), N-acetylated, reduced with NaBD$_4$ and acetylated with a 1:1 pyridine-acetic anhydride mixture (100 °C, 30 min). To release acidic sugar components, LPSs and the OPS were subjected to methanolysis (1 M HCl in methanol, 85 °C, 16 h), carboxyl reduction with NaBD$_4$ in aqueous 50% methanol, hydrolysis with 2 M CF$_3$CO$_2$H and acetylation. The products were identified as alditol acetates by GC-MS [51] on a Hewlett-Packard HP5890A-HP5971 instrument equipped with an HP-5ms (SLB-5ms) capillary column (30 m × 0.25 mm; Supelco, St. Louis, MO, USA), applying a temperature gradient of 150 °C (5 min) to 310 °C at 5 °C min$^{-1}$.

The absolute configuration of monosaccharides was determined by GC of acetylated (S)-2-butyl glycosides using authentic sugars as standards [36].

Methylation of the OPS was performed by the procedure of Hakomori [52]. The permethylated OPS was subjected to hydrolysis in 2 M CF$_3$CO$_2$H (120 °C, 2 h), N-acetylation, and reduction with NaBD$_4$. Partially methylated alditols (PMAA) were converted into acetate derivatives, and analyzed by GC-MS as above.

For fatty acid analysis, a sample of the lipid A (1 mg) was subjected to methanolysis in 2 M methanolic HCl (85 °C, 12 h). The resulting fatty acid methyl esters were extracted with hexane and converted to their O-trimethylsilyl (O-TMS) derivatives, as described [50,53]. The methanol layer, containing the methyl glycosides was dried and acetylated with pyridine-acetic anhydride mixture. The fatty acid derivatives, as well as acetylated methyl glycosides were analyzed by GC-MS as above.

3.4. NMR Spectroscopy

1D $^1$H NMR, $^{13}$C NMR and 2D NMR experiments were recorded in a D$_2$O solution at 32 °C using a Bruker Avance III 700 MHz spectrometer (operating frequencies 700.43 MHz for $^1$H NMR and 176.14 MHz for $^{13}$C NMR) and applying standard Bruker software (Bruker, TopSpi, Rheinstetten, Germany). Chemical shifts were reported relative to external acetone as reference (δ$_H$ 2.225, δ$_C$ 31.45). The following homo- and heteronuclear correlated two-dimensional spectra were used for general assignments: $^1$H, $^1$H DQF-COSY, TOCSY, ROESY, $^1$H,$^{13}$C HSQC, and $^1$H,$^{13}$C HMBC.

3.5. Mass Spectrometry Analysis

ESI FT-ICR MS was performed in negative ion mode using a hybrid Apex Qe FT-ICR MS instrument (Bruker Daltonics), equipped with a 7 Tesla actively shielded magnet and an Apollo dual ion source. Samples (~10 ng μL$^{-1}$) were sprayed at a flow rate of 2 μL min$^{-1}$. Capillary entrance voltage was set to 3.8 kV, and dry gas temperature to 200 °C. For unspecific fragmentation the DC offset (collision voltage) of the collision cell was set from 5 V to 30 V. Under these conditions the labile linkage between the lipid A and the core oligosaccharide is cleaved [54,55] resulting in intensive
Y− and B− fragments representing the lipid A and the core oligosaccharide moieties (according to the nomenclature of Domon and Costello [33]). The mass spectra were charge deconvoluted and mass values given in all spectra and tables refer to the monoisotopic signal of the neutral molecules. Mass calibration was done externally by well-characterized LPS of known structure.

3.6. SDS-PAGE

LPS preparations were separated in 12.5% SDS-Tricine polyacrylamide electrophoresis gel and bands were visualized by silver staining after oxidation with periodate [56].

4. Conclusions

Aeromonas sp. bacteria are common inhabitants of aquatic environments and have been described in relation to fish and human diseases, particularly food-borne associated [2,18]. The cell envelope of Aeromonas, as that of other Gram-negative bacteria, contains LPS, a crucial amphiphilic and immunodominant constituent of the outer membrane [57]. The OPS of the LPS is one of the most variable components on the cell surface, providing the basis for serotyping of bacteria. As it was shown that specific Aeromonas, as well as Cronobacter O serotypes, were associated with enteritis epidemics [5,58], especially in children, it is important to identify epidemiologically relevant strains and to understand the immunochemical aspects of antigen specificity within the serogroups.

While most OPS are distinct among different genera, the core region of LPS, in particular the inner part, tends to be conserved within a genus or even family. The fact that the inner core of LPS from distantly related bacteria shares structural features is a reflection of evolutionary relationship and the importance of this region in outer membrane integrity [57]. On the other hand, the outer core shows more structural diversity, as might be expected for a region exposed to the selective pressures of host responses, location of bacteriophage receptors, and environmental stress. Recently, the complete core structure of LPS from A. hydrophila strain AH-901, which is a mutant in a gene encoding mannose transferase, has been published. The core nonasaccharide was composed of two D,D-Hep residues in combination with four L,D-Hep. Two other sugars, i.e., β-glucose and α-glucosamine, as well as one residue of α-3-deoxy-d-manno-oct-2-ulosonic acid, at the reducing end, were also detected. Additionally, one of D,D-heptose residue was non-stoichiometrically substituted with β-galactopyranose. No charged groups were reported except for one phosphate group at the 4-position of the Kdo unit [59,60].

Moreover, our latest findings revealed that the core oligosaccharide with the prevalence of heptose residues and the composition Hep6Hex1HexN1Kdo1P1 represents a structure shared by LPS core part of the strains belonging to the species A. hydrophila and A. bestiarum [27,59,60].

Interestingly, our present results show that the core region may vary to some extent within Aeromonas spp. bacteria. In this work, water- and phenol-soluble LPS preparations isolated from A. veronii strain Bs19 have been structurally characterized. Compositional analysis identified D,D-Hep and L,D-Hep, and ESI MS experiments confirmed that the core decasaccharide had a different structure than those established earlier [27,59,60], namely Hep3Hex3HexN1Kdo1P1. Some differences were also noticed in the composition and acylation pattern of the lipid A. Although LPS glycoforms had tetra-acylated and hexa-acylated lipid A species with amide-linked 14:0(3-OH) and a backbone comprising a bisphosphorylated GlcN disaccharide, an AraN residue was exclusively detected as a
non-stoichiometric substituent in the lipid A from LPS of the phenol-soluble fraction. Additionally, some lipid A species contained ester-linked 12:0 and 14:0.

In the future, we would like to focus on structural studies of LPS heterogeneity within the strains belonging to the group *Aeromonas sobria* complex and combine the data with the elucidation of the location and organization of LPS gene clusters.

**Acknowledgments**

This work was financially supported by the grant from the National Science Centre (Decision No. DEC-2011/03/B/NZ1/01203). The authors wish to thank Brigitte Kunz for recording ESI FT-ICR mass spectra and Buko Lindner for his valuable discussion and constructive suggestions during elaborating of the mass spectrometry data. We also wish to thank Heiko Käßner for recording NMR spectra and Hermann Moll for his interest and assistance in chemical analyses (both from Research Center Borstel, Borstel, Germany). The authors especially thank Hubert Pietras for his technical assistance (Department of Genetics and Microbiology, Maria Curie-Skłodowska University, Lublin, Poland).

**Conflicts of Interest**

The authors declare no conflict of interest.

**References**

1. Janda, J.M.; Duffy, P.S. Mesophilic aeromonads in human diseases: current taxonomy, laboratory infection and infectious diseases spectrum. *Rev. Infect. Dis.* **1988**, *10*, 980–997.
2. Janda, J.M. Recent advances in the study of the taxonomy, pathogenicity and infectious syndromes with the genus *Aeromonas*. *Clin. Microbiol. Rev.* **1991**, *4*, 397–410.
3. Nawaz, M.; Khan, S.A.; Khan, A.A.; Sung, K.; Tran, Q.; Kerdahi, K.; Steele, R. Detection and characterization of virulence genes and integrons in *Aeromonas veronii* isolated from catfish. *Food Microbiol.* **2010**, *27*, 327–331.
4. Araujo, R.M.; Arribas, R.M.; Pares, R. Distribution of *Aeromonas* species in waters with different levels of pollution. *J. Appl. Bacteriol.* **1991**, *71*, 182–186.
5. Janda, J.M.; Abbott, S. The genus *Aeromonas*: taxonomy, pathogenicity, and infection. *Clin. Microbiol. Rev.* **2010**, *23*, 35–73.
6. Rahman, M.; Colque-Navarro, P.; Kühn, I.; Huys, G.; Swings, J.; Möllby, R. Identification and characterization of pathogenic *Aeromonas veronii* bv. *sobria* associated with epizootic ulcerative syndrome in fish in Bangladesh. *Appl. Environ. Microbiol.* **2002**, *68*, 650–655.
7. Cai, S.-H.; Wu, Z.-H.; Jian, J.-C.; Lu, Y.-S.; Tang, J.F. Characterization of pathogenic *Aeromonas veronii* bv. *veronii* associated with ulcerative syndrome from Chinese longsnout catfish (*Leiocassis longirostris* Günther). *Braz. J. Microbiol.* **2012**, *43*, 382–388.
8. Holmberg, S.D.; Schell, W.L.; Fanning, G.R.; Wachsmuth, I K.; Blake, P.A.; Brenner, D.J.; Farmer, J.J. *Aeromonas* intestinal infections in the United States. *Ann. Intern. Med.* **1986**, *105*, 683–689.
9. Ali, A.; Carnahan, A.M.; Altwegg, M.; Luthy-Hottenstein, J.; Joseph, S.W. *Aeromonas bestiarum* sp. nov. (formerly genomospecies DNA group 2 *A. hydrophila*), a new species isolated from non human sources. *Med. Microbiol. Lett.* **1996**, *5*, 156–165.
10. Kahajanchi, B.K.; Fadl, A.A.; Borchardt, M.A.; Berg, R.L.; Horneman, A.J.; Stemper, M.E.; Joseph, S.W.; Moyer, N.P.; Sha, J.; Chopra, A.K. Distribution of virulence factors and molecular fingerprinting of *Aeromonas* species isolates from water and clinical samples: suggestive evidence of water-to-human transmission. *Appl. Environ. Microbiol.* **2010**, *76*, 2313–2325.
11. Figueras, M.J. Clinical relevance of *Aeromonas* spp. *Rev. Clin. Microbiol.* **2005**, *16*, 145–153.
12. Martinez-Murcia, A.J.; Borrell, N.; Figueras, M.J. Typing of clinical and environmental *Aeromonas veronii* strains based on the 16S-23S rDNA spacers. *FEMS Immunol. Med. Microbiol.* **2000**, *28*, 225–232.
13. Dooley, J.S.G.; Lallier, R.; Shaw, D.H.; Trust, T.J. Electrophoretic and immunochemical analyses of the lipopolysaccharides from various strains of *Aeromonas hydrophila*. *J. Bacteriol.* **1985**, *164*, 263–269.
14. Merino, S.; Rubires, X.; Aguillar, A.; Guillot, J.F.; Tomas, J.M. The role of the O-antigen lipopolysaccharide on the colonization *in vivo* of the germfree chicken gut by *Aeromonas hydrophila* serogroup O:34. *Microb Pathog.* **1996**, *20*, 325–333.
15. Aguilar, A.; Merino, S.; Rubires, X.; Tomas, J. Influence of osmolarity on lipopolysaccharides and virulence of *Aeromonas hydrophila* serotype O:34 strains grown at 37 degrees C. *Infect. Immun.* **1997**, *65*, 1245–1250.
16. Rabaan, A.A.; Gryllos, I.; Tomas, J.M.; Shaw, J.G. Motility and polar flagellum are required for *Aeromonas caviae* adherence to HEp-2 cells. *Infect. Immun.* **2001**, *69*, 4257–4267.
17. Garduno, R.A.; Moore, A.R.; Oliver, G.; Lizama, A.L.; Garduno, E.; Kay, W.W. Host cell invasion and intracellular resistance by *Aeromonas salmonicida*: role of the S-layer. *J. Clin. Microbiol.* **2000**, *46*, 660–668.
18. Sakazaki, R.; Shimada, T. O-serogrouping for mesophilic *Aeromonas* strains. *Jpn. J. Med. Sci.* **1984**, *37*, 247–255.
19. Nandapalan, N.; Chang, B.J. Production and characterization of monoclonal antibodies to *Aeromonas sobria* surface antigens. *FEMS Microbiol. Immunol.* **1989**, *47*, 515–524.
20. Francki, K.T.; Chang, B.J.; Mee, B.J.; Collignon, P.J.; Susai, V.; Keese, P.K. Identification of genes associated with copper tolerance in an adhesion-defective mutant of *Aeromonas veronii* biovar *sobria*. *FEMS Immunol. Med. Microbiol.* **2000**, *29*, 115–121.
21. Hickman-Brenner, F.W.; MacDonald, K.L.; Steigerwalt, A.G.; Fanning, G.R.; Brenner, D.J.; Farmer, J.J., III. *Aeromonas veronii*, a new ornithine decarboxylase-positive species that may cause diarrhea. *J. Clin. Microbiol.* **1987**, *25*, 900–906.
22. Mencacci, A.; Cenci, E.; Mazzolla, R.; Farinella, S.; D’Alò, F.; Vitali, M.; Bistoni, F. *Aeromonas veronii* biovar *veronii* septicemia and acute suppurative cholangitis in a patient with hepatitis B. *J. Med. Microbiol.* **2003**, *52*, 727–730.
23. Roberts, M.T.M.; Enoch, D.A.; Harris, K.A.; Karas, J.A. *Aeromonas veronii* biovar *sobria* bacteraemia with septic arthritis confirmed by 16S rDNA PCR in an immunocompetent adult. *J. Med. Microbiol.* **2006**, *55*, 241–243.
24. Wang, Z.; Liu, X.; Dacanay, A.; Harrison, B.A.; Fast, M.; Colquhoun, D.J.; Lund, V.; Brown, L.L.; Li, J.; Altman, E. Carbohydrate analysis and serological classification of typical and atypical isolates of *Aeromonas salmonicida*: A rationale for the lipopolysaccharide-based classification of *A. salmonicida*. *Fish Shellfish Immun.* 2007, 23, 1095–1106.

25. Knirel, Y.A.; Shashkov, A.S.; Senchenkova, S.N.; Merino, S.; Tomas, J.M. Structure of the O-specific polysaccharide of *Aeromonas hydrophila* O:34; a case of random O-acetylation of 6-deoxy-L-talose. *Carbohydr. Res.* 2002, 337, 1381–1386.

26. Nazarenko, E.L.; Crawford, R.J.; Iwanowa, E.P. The structural diversity of carbohydrate antigens of selected Gram-negative marine bacteria. *Mar. Drugs* 2011, 9, 1914–1954.

27. Turska-Szewczuk, A.; Lindner, B.; Komaniecka, I.; Kozinska, A.; Pekala, A.; Choma, A.; Holst, O. Structural and immunochemical studies of the lipopolysaccharide from the fish pathogen, *Aeromonas bestiarum* strain K296, serotype O18. *Mar. Drugs* 2013, 11, 1235–1255.

28. Kozinska, A.; Pekala, A. Serotyping of *Aeromonas* species isolated from Polish fish farms in relation to species and virulence phenotype of the bacteria. *Bull. Vet. Inst. Pulawy* 2010, 54, 315–320.

29. Turska-Szewczuk, A.; Kozinska, A.; Russa, R.; Holst, O. The structure of the O-specific polysaccharide from the lipopolysaccharide of *Aeromonas bestiarum* strain 207. *Carbohydr. Res.* 2010, 345, 680–684.

30. Turska-Szewczuk, A.; Guz, L.; Lindner, B.; Pietras, H.; Russa, R.; Holst, O. Structural characterization of the O-specific polysaccharide from the lipopolysaccharide of fish pathogen *Aeromonas bestiarum* strain P1S. *Carbohydr. Res.* 2011, 346, 815–821.

31. Kozinska, A.; Figueras, M.J.; Chacon, M.R.; Soler, L. Phenotypic characteristics of *Aeromonas* genomospecies isolated from common carp (*Cyprinus carpio* L.). *J. Appl. Microbiol.* 2002, 93, 1034–1041.

32. Westphal, O.; Jann, K. Bacterial lipopolysaccharide. Extraction with phenol-water and further applications of the procedure. *Meth. Carbohydr. Chem.* 1965, 5, 83–91.

33. Domon, B.; Costello, C.E. A systematic nomenclature for carbohydrate fragmentations in FAB MS/MS spectra of glycoconjugates. *Glycoconj. J.* 1988, 5, 397–409.

34. MacLean, L.L.; Perry, M.B. Structural characterization of the serotype O:5 O-polysaccharide antigen of the lipopolysaccharide of *Escherichia coli* O:5. *Biochem. Cell Biol.* 1997, 75, 199–205.

35. Turska-Szewczuk, A.; Lindner, B.; Pekala, A.; Palusinska-Szysz, M.; Choma, A.; Russa, R.; Holst, O. Structural analysis of the O-specific polysaccharide from the lipopolysaccharide of *Aeromonas veronii* bv. *sobria* strain K49. *Carbohydr. Res.* 2012, 353, 62–68.

36. Leontein, K.; Lindberg, B.; Lönngren, J. Assignment of absolute configuration of sugars by GLC of their acetylated glycosides formed from chiral alcohols. *Carbohydr. Res.* 1978, 62, 359–362.

37. Lipkind, G.M.; Shashkov, A.S.; Knirel, Y.A.; Vinogradov, E.V.; Kochetkov, N.K. A computer-assisted structural analysis of regular polysaccharides on the basis of 13C-n.m.r. data. *Carbohydr. Res.* 1988, 175, 59–75.

38. Shashkov, A.S.; Vinogradov, E.V.; Knirel, Y.A.; Nifant’ev, N.E.; Kochetkov, N.K.; Dabrowski, J.; Kholodkova, E.V.; Stanislavsky, E.S. Structure of the O-specific polysaccharide of *Salmonella arizonae* O45. *Carbohydr. Res.* 1993, 241, 177–188.
39. Shashkov, A.S.; Paramonov, N.A.; Veremeychenko, S.P.; Grosskurth, H.; Zdorovenko, G.M.; Knirel, Y.A.; Kochetkov, N.K. Somatic antigens of pseudomonads: structure of the O-specific polysaccharide of *Pseudomonas fluorescens* biovar B, strain IMV 247. *Carbohydr. Res.* **1998**, *306*, 297–303.

40. Senchenkova, S.N.; Shashkov, A.S.; Laux, P.; Knirel, Y.A.; Rudolph, K. The O-chain polysaccharide of *Xanthomonas campestris* pv. *begoniae* GSPB 525 is a partially L-xylosylated rhamnan. *Carbohydr. Res.* **1999**, *319*, 148–153.

41. Carillo, S.; Silipo, A.; Perino, V.; Lanzetta, R.; Parrilli, M.; Molinaro, A. The structure of the O-specific polysaccharide from the lipopolysaccharide of *Burkholderia anthina*. *Carbohydr. Res.* **2009**, *344*, 1697–1700.

42. Katzenellenbogen, E.; Romanowska, E.; Kocharova, N.A.; Knirel, Y.A.; Shashkov, A.S.; Kochetkov, NK. Structure of a glycerol teichoic acid-like O-specific polysaccharide of *Hafnia alvei* 1205. *Carbohydr. Res.* **1992**, *231*, 249–260.

43. Silipo, A.; Leone, S.; Lanzetta, R.; Parrilli, M.; Sturiale, L.; Garozzo, D.; Nazarenko, E.L.; Gorshkova, R.P.; Ivanova, E.P.; Gorshkova, N.M.; Molinaro, A. The complete structure of the lipooligosaccharide from the halophilic bacterium *Pseudoalteromonas issachenkonii* KMM 3549. *Carbohydr. Res.* **2004**, *339*, 1985–1993.

44. Jansson, P.E.; Kenne, L.; Widmalm, G. Computer-assisted structural analysis of polysaccharides with an extended version of CASPER using ¹H- and ¹³C-NMR data. *Carbohydr. Res.* **1989**, *188*, 169–191.

45. Toukach, F.V.; Bartodziejska, B.; Senchenkova, S.N.; Wykrota, M.; Shashkov, A.S.; Rozalski, A.; Knirel, Y.A. Structure of a new acidic O-antigen of *Proteus vulgaris* O22 containing O-acetylated 3-acetamido-3,6-dideoxy-D-glucose. *Carbohydr. Res.* **1999**, *318*, 146–153.

46. Vinogradov, E.V.; Petersen, B.O.; Thomas-Oates, J.E.; Duus, J.O.; Brade, H.; Holst, O. Characterization of a novel branched tetrasaccharide of 3-deoxy-D-manno-oct-2-ulopyranosonic acid. The structure of the carbohydrate backbone of the lipopolysaccharide from *Acinetobacter baumannii* strain NCTC 10303 (ATCC 17904). *J. Biol. Chem.* **1998**, *273*, 28122–28131.

47. Kumirska, J.; Szafranek, J.; Czerwicka, M.; Paszkiewicz, M.; Dziadziuszko, H.; Kunikowska, D.; Stepnowski, P. The structure of the O-specific polysaccharide isolated from the lipopolysaccharide of *Salmonella* Dakar (serogroup O:28). *Carbohydr. Res.* **2007**, *342*, 2138–2143.

48. MacLean, L.L.; Vinogradov, E.; Perry, M.B. The structure of the antigenic O-polysaccharide in the lipopolysaccharide of enterohaemorrhagic *Escherichia coli* serotype O71:H12. *Biochem. Cell. Biol.* **2010**, *88*, 439–444.

49. Dziadziuszko, H.; Kumirska, J.; Muža, S.; Czerwicka, M.; Lubecka, E.A.; Stepnowski, P.; Kunikowska, D. Immunochemical studies of *Salmonella* Dakar and *Salmonella* Telaviv O-antigens (serogroup O:28). *FEMS Microbiol. Lett.* **2012**, *326*, 55–61.

50. Komaniecka, I.; Choma, A.; Lindner, B.; Holst, O. The structure of a novel lipid A from the lipopolysaccharide of *Bradyrhizobium elkanii* containing three mannose units in the backbone. *Chem. Eur. J.* **2010**, *16*, 2922–2929.
51. Russa, R.; Urbanik-Sypniewska, T.; Lindström, K.; Mayer, H. Chemical characterization of two lipopolysaccharide species isolated from Rhizobium loti NZP2213. Arch. Microbiol. 1995, 163, 345–351.

52. Hakomori, S. A rapid permethylation of glycolipid and polysaccharide catalyzed by methylsulfinyl carbanion in dimethyl sulfoxide. J. Biochem. 1964, 55, 205–208.

53. Pieretti, G.; Corsaro, M.M.; Lanzetta, R.; Parrilli, M.; Vilches, S.; Merino, S.; Tomas, J.M. Structure of the core region from the lipopolysaccharide of Plesiomonas shigelloides strain 302-73 (serotype O1). Eur. J. Org. Chem. 2009, 2009, 1365–1371.

54. Kondakova, A.; Lindner, B. Structural characterization of complex bacterial glycolipids by Fourier transform ion cyclotron mass spectrometry. Eur. J. Mass Spectrom. 2005, 11, 535–546.

55. Klein, G.; Lindner, B.; Brabetz, W.; Brade, H.; Raina, S. Escherichia coli K-12 suppressor-free mutants lacking early glycosyltransferases and late acyltransferases: minimal lipopolysaccharide structure and induction of envelope stress response. J. Biol. Chem. 2009, 284, 15369–15389.

56. Turska-Szewczuk, A.; Palusinska-Szysz, M.; Russa, R. Structural studies of O-polysaccharide chain from the lipopolysaccharide of symbiotically enhanced mutant Mlo-13 of Mesorhizobium loti NZP2213. Carbohydr. Res. 2008, 343, 477–482.

57. Raetz, C.R.H.; Whitfield, C. Lipopolysaccharide endotoxins. Annu. Rev. Biochem. 2002, 71, 635–700.

58. Sun, Y.; Wang, M.; Wang, Q.; Cao, B.; He, X.; Li, K.; Feng, L.; Wang, L. Genetic analysis of the Cronobacter sakazakii O4 to O7 O-antigen gene clusters and development of a PCR assay for identification of all C. sakazakii O serotypes. Appl. Environ. Microbiol. 2012, 78, 3966–3974.

59. Knirel, Y.A.; Vinogradov, E.; Jimenez, N.; Merino, S.; Tomas, J.M. Structural studies on the R-type lipopolysaccharide of Aeromonas hydrophila. Carbohydr. Res. 2004, 339, 787–793.

60. Jimenez, N.; Canals, R.; Lacasta, A.; Kondakova, A.; Lindner, B.; Knirel, Y.A.; Merino, S.; Regue, M.; Tomas, J.M. Molecular analysis of three Aeromonas hydrophila AH-3 (Serotype O34) lipopolysaccharide core biosynthesis gene clusters. J. Bacteriol. 2008, 190, 3176–3184.

© 2014 by the authors; licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution license (http://creativecommons.org/licenses/by/3.0/).