

1 Pyrolysis of end-of-life polystyrene in a 2 pilot-scale reactor: maximizing styrene 3 production

4 Azd Zayoud^{1,3}, Hang Dao Thi¹, Marvin Kusenber¹, Andreas Eschenbacher¹, Uros Kresovic²,
5 Nick Alderweireldt², Marko Djokic¹, Kevin M. Van Geem¹

6 ¹Laboratory for Chemical Technology, Department of Materials, Textiles and Chemical
7 Engineering, Ghent University, Gent 9052, Belgium;

8 ²Indaver N.V. Belgium, Mechelen 2800, Belgium;

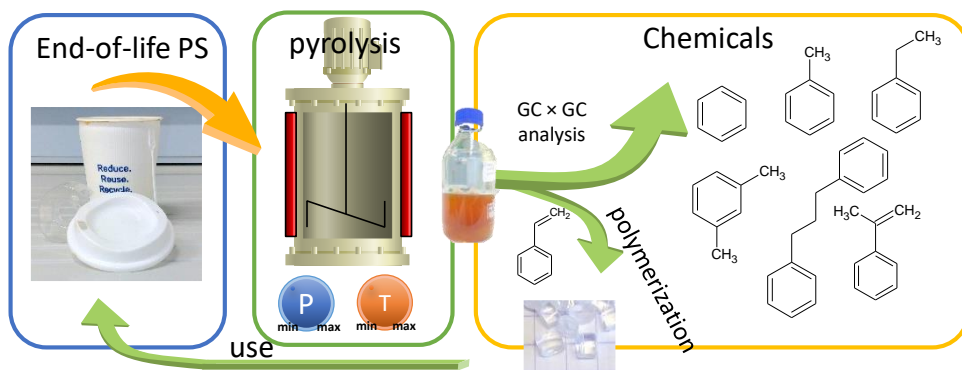
9 ³Université Catholique de Louvain, Institute of Mechanics, Materials and Civil Engineering,
10 1348 Louvain-la-Neuve, Belgium

11 Highlights

- 12 • End-of-life polystyrene was pyrolyzed in CSTR under vacuum and atmospheric
13 pressure
- 14 • Vacuum reduced the yield of styrene oligomers compared to atmospheric pressure
- 15 • Maximum styrene yield was 55.9 wt.% at 550 °C and 0.02 bara operating conditions
- 16 • Styrene dimers, trimers and tetramers were in detail quantified by GC × GC-FID/-MS

17

18 Graphical abstract



19

20

21 Abstract

22 Chemical recycling of polystyrene (PS) via pyrolysis is of great industrial, and academic
23 interest, with styrene being the primary product of interest. To identify the optimal process
24 conditions, the pyrolysis of end-of-life PS was studied in a pilot-scale unit consisting of an
25 extruder, and a continuous stirred tank reactor (CSTR). The PS was pyrolyzed with continuous
26 feeding at a pressure range from 0.02 to 1.0 bara, and a temperature range from 450 to
27 600 °C, giving primarily styrene, other mono-aromatics, and oligomers. The comprehensive
28 two-dimensional gas chromatography (GC × GC) coupled with flame ionization detector (FID),
29 and time-of-flight mass spectrometer (ToF-MS) as well as GC with thermal conductivity
30 detector (TCD) were used to characterize the liquid, and gaseous products exhaustively.

31 The styrene yield increased from 36 wt.% at 1.0 bara, and 450 °C to 56 wt.% at 0.02 bara, and
32 550 °C. Working under a vacuum enhanced the styrene recovery at all corresponding
33 temperature levels. The yield of benzene, toluene, ethylbenzene, and xylene (BTEX) increased
34 from 4 wt.% at 450 °C, and 0.02 bara to 17 wt.% at 450 °C, and 1.0 bara. The experimental
35 results have been used in a mathematical model that can explain the combined effect of
36 temperature, and pressure on the yield of the primary products. The present work illustrates
37 the potential of a continuous pyrolysis process for end-of-life PS, and paves the way for this
38 technology to be rapidly transferred from mere laboratory use to industrial processes in the
39 circular (petro-) chemical industry.

40 **Keywords:** pyrolysis, waste plastic, continuous process, polystyrene, CSTR, mathematical
41 optimization

42 1 Introduction

43 The steadily increasing plastics production due to a continuously growing global demand
44 exacerbates the end-of-life plastics problem. In 2018, the world production exceeded 350
45 million metric tons, where the European production share was around 17% (Garside, 2019).
46 The intensive use of plastics has put the spotlight on how to resolve the end-of-life plastic
47 problem (PlasticsEurope, 2018). Additionally, China, and India banned the import of end-of-
48 life plastics (Brooks et al., 2018). As a result, treating end-of-life plastics domestically becomes
49 unavoidable for many countries. In Europe, the target is set to reuse, and recycle 60 % of all
50 plastic packaging by 2030, and 100% of all plastic packaging will be either reused, recycled, or
51 recovered by 2040 (PlasticsEurope, 2019a).

52 PS is one of the main demanded polymers, and subsequently one of the major end-of-life
53 plastic types (PlasticsEurope, 2019b). And PS pyrolysis has been studied intensively during the
54 past decades. However, to date, the reported studies in open literature have been focused on
55 processing virgin PS and ultra-pure waste PS but not (contaminated) end-of-life PS. Madorsky
56 and Straus were one of the first to perform PS pyrolysis experiments in a laboratory-scale
57 batch reactor at vacuum pressure and a temperature range from 340 to 420 °C; these
58 researchers reported a maximum styrene recovery of 42 wt.% at 420 °C (Madorsky and Straus,
59 1948). In 1981, Ogino and Nagy used a micro-pyrolyzer to process PS at vacuum pressure and
60 a temperature range between 400 and 500 °C, achieving a high styrene yield of 84.5 wt.%
61 (Ogino and Nagy, 1981). Using a laboratory-scale fixed bed reactor, Achilias et al. pyrolyzed PS
62 at 510 °C and obtained a styrene selectivity of 63.9 wt.% in their condensed liquid product,
63 which implies an overall styrene yield of 58.7 wt.% (Achilias et al., 2007). Mo et al. applied a
64 response surface method (RSM) to maximize the styrene recovery using a laboratory-scale

65 semi-batch horizontal tube furnace at atmospheric pressure and found a maximum styrene
66 yield of 60.9 wt.% at 490 °C (Mo et al., 2014). A microwave laboratory-scale reactor was used
67 to pyrolyze PS in a temperature range from 464 to 678°C, and reported a styrene yield of 66.0
68 wt.% (Undri et al., 2014). Similarly, Bartoli et al. performed PS pyrolysis with a microwave
69 laboratory-scale reactor at vacuum pressure and a temperature range of 301-536 °C and
70 reported a styrene yield of 60.6 wt.% (Bartoli et al., 2015). The earliest works by Kim et al.
71 using laboratory-scale reactors proved the concept of recovering monomers of PS through a
72 pyrolysis process (Kim et al., 1999). However, most of the previous works used either batch or
73 manual feeding modes (Chauhan et al., 2008; Liu et al., 2000; Park et al., 2003; Williams and
74 Williams, 2010). The batch and manual feeding modes have inherent downtime, high
75 operating cost, and product variability, as a result, these batch processes and manual feeding
76 reactors have little potential to be scaled up (Qureshi et al., 2018). Anyhow, these preliminary
77 works have paved the way for advancing the experimental end-of-life plastics pyrolysis works.
78 Further investigation of the PS pyrolysis process with continuous feeding is required since only
79 a few PS pyrolysis experiment works were performed in continuous mode (Ando et al., 1974;
80 Kaminsky et al., 2004; Park et al., 2020).

81 Even though data is scarce, PS pyrolysis gains momentum on an industrial scale, with several
82 demonstration projects being announced (PolystyreneLoop, 2021; Pyrolyze.B.V., 2021;
83 Smalley, 2019; Victory, 2020). Until today, PS pyrolysis in a CSTR with continuous feeding has
84 not been implemented on an industrial scale yet. Instead, reactors such as a twin-screw and a
85 microwave reactor have been used to process PS (Agilyx, 2020; Doucet, 2020; Doucet et al.,
86 2016; Qureshi et al., 2019). According to classical chemical engineering principles, scaling up
87 step by step of the continuous processing capacity is considered best practice. Therefore,

88 further investigations on PS pyrolysis in a CSTR with continuous feeding are advised before
89 developing and building an industrial size plant. The CSTR is a robust reactor type, offering the
90 flexibility of operating at different temperatures and pressure levels, *viz.* vacuum, atmospheric
91 pressure and elevated pressure. Based on this flexibility in operating conditions, this reactor
92 type was selected for the present work. An additional advantage is that it facilitates the
93 processing of a broad range of plastics types such as PE, PP, PS, and mixed plastic waste.

94 Besides the lack of studies on PS pyrolysis in continuous feeding mode and operating under
95 vacuum, there is limited knowledge about the detailed composition of the oligomers of
96 styrene formed during the pyrolysis of PS. With more advanced analytical techniques, it is
97 possible to gain new insights into PS pyrolysis and the impact of the process conditions on
98 monomer and oligomer formation. In particular, under vacuum, it is expected that more
99 styrene can be formed because secondary reactions are minimized. In the open literature, the
100 PS pyrolyzate analysis was only carried out using one-dimensional gas chromatography (1D-
101 GC) with typically either a thermal conductivity detector (TCD), a flame ionization
102 detector (FID), or a time of flight mass spectrometry detector (TOF-MS) (Kaminsky and Franck,
103 1991; Kim et al., 1999; Ogino and Nagy, 1981; Park et al., 2003; Park et al., 2020; Williams and
104 Williams, 2010). In the characterized pyrolyzate, the unidentified compounds ranged between
105 7 and 18 wt.%, possibly because of insufficient identification with databases and limitations
106 of the sampling system (Artetxe et al., 2015; Kaminsky et al., 2004; Park et al., 2020). Kaminsky
107 et al. identified 12 PS pyrolyzate compounds (Kaminsky, 2021; Kaminsky et al., 2004) and Undri
108 et al. quantified the compounds with a concentration $\geq 0.2\%$ (Undri et al., 2014).
109 Comprehensive two-dimensional gas chromatography (GC \times GC) (Dalluge et al., 2002) can be
110 beneficial for the quantitative analysis because of its intrinsic higher sensitivity (Dijkmans et

111 [al., 2015; Phillips and Beens, 1999](#)). The most advanced study -for detailed compositional
112 characterization of plastic waste pyrolysis oil- considered GC × GC coupled to multiple
113 detectors such as FID, TOF-MS, a sulfur chemiluminescence detector (SCD), and nitrogen
114 chemiluminescence detector (NCD), enabling the quantification of impurities ([Toraman et al.,](#)
115 [2014](#)).

116 The primary aim of the present study was to improve the understanding of the effect of
117 temperature and pressure on the continuous pyrolysis process of end-of-life PS feedstock
118 which was obtained from industrial scale sorting plant. This was performed in a CSTR pilot-
119 scale continuous feeding unit. Particular attention was paid to the detailed quantification and
120 identification of the formed oligomers, the pyrolyzate was analyzed in detail using the state
121 of the art comprehensive two dimensional GC × GC-FID/-MS. Experiments were conducted at
122 three pressure levels (0.02, 0.5, and 1.0 bara), and four temperature levels (450, 500, 550, and
123 600 °C); interestingly, the vacuum pressure was found to have an insignificant effect on the
124 pyrolyzate yield, but a noteworthy effect on the composition of the pyrolyzate products. The
125 maximum styrene and maximum liquid pyrolyzate yields were obtained at vacuum pressure
126 (0.02 bara) and an operating temperature of ~550 °C. These findings can be used for
127 optimizing the operating conditions of an industrial plant. The obtained products were
128 analyzed using a comprehensive set of analytical techniques. The obtained experimental
129 results were interpolated using a mathematical model to predict the optimal operating
130 conditions.

131 2 Materials and Methods

132 2.1 Materials

133 End-of-Life PS (Coolrec, Belgium) was used as a feedstock for the pyrolysis experiments. The
134 end-of-life PS was collected, washed, shredded, and granulated. The composition of the used
135 waste fractions has been estimated (Roosen et al., 2020a), and is given by 94.2 % (E)PS, 0.2 %
136 PET, 1.2 % PE, 0.5 % EVOH, 0.5 % PA, 0.3 % PUR, and 3.0 % paper; and the weight-based
137 elemental composition (CHNS/O) of the end-of-life PS is given by 86.6 % C, 8.2 % H, 0.5 % N,
138 <level of detection (LOD) S, 2.7% O and 2.0 % metal and halogen contents (Roosen et al.,
139 2020a). Additionally, the ultimate analysis was carried out for pyrolyzate products [e.g. (Exp.
140 4)], the measured CHNS/O is given by 86.3 % C, 7.7 % H, 0.6 % N, <LOD S, 5.3 % O and other
141 compounds. Carbon disulfide, with a purity of 99.9% (Sigma-Aldrich, Belgium), was used as a
142 solvent to dissolve the PS pyrolyzate before the GC × GC analysis. 3-Chlorothiophene,
143 procured with a purity of 97% (Sigma-Aldrich, Belgium), was used as an internal standard in
144 GC × GC analysis. Benzene, toluene, and styrene with a purity of ≥99% (Acros Organics,
145 Belgium) were used for external calibration. Analytical gasses (nitrogen, helium, oxygen, and
146 hydrogen) used for GC × GC had a minimum purity of 99.999% (AirLiquide, Belgium). The 4-
147 tert-Butylcatechol inhibitor with a purity of ≥99.0% (Sigma-Aldrich, Belgium) was added to the
148 pyrolyzate samples to avoid auto-polymerization.

149 2.2 Experimental apparatus and procedure

150 2.2.1 Pyrolysis pilot unit

151 The PS pyrolysis experiments were performed using an in-house developed continuous
152 pyrolysis pilot-scale unit at the Laboratory for Chemical Technology (LCT, Ghent University,

153 Ghent 9052, Belgium). As shown in [Figure 1](#), the pyrolysis unit consists of three main sections.

154 In the feeding section, a LabTech single screw extruder (1) (Model: LE25-30/CV, Thailand) with

155 a feeding rate range of $0.1-10 \text{ kg}\cdot\text{h}^{-1}$, and four dedicated heating zones are used to pre-heat

156 and melt the granulated plastic feedstock and to feed it to the reactor. The extruder's outlet

157 is connected to the reactor's inlet by a heated transfer line; more information on the

158 extruder's specifications can be found in the ([supplementary information](#)).

159 In the reaction section, a Parr reactor (Model: 4584, U.S.A) is used for pyrolysis. The reactor

160 has a volume of 5.7 L (1.5 gallons) and the heat is provided using a heating jacket. The reactor

161 conditions are controlled through a PC (3) using SpecView software and a PID controller. The

162 reactor is also equipped with three K-type thermocouples, a stirrer, shaft magnetic coupling,

163 a manometer, and a pressure transducer. A nitrogen bottle of $\geq 99.99\%$ purity (4) (AirLiquide,

164 Belgium) is connected to the reactor for purging before the experiment and the nitrogen flow

165 is controlled through a volume flow controller (5) (KROHNE, MD: 2018, Germany).

166 In the condensation section, three condensers (6) operated in series captured the pyrolyzate

167 and the temperature of these condensers is controlled by using an external LAUDA cooler (7)

168 (RE 420 G, Germany). Each condenser set consists of three valves: **a**) valves ([Figure 1](#), v-201,

169 v-188, and v-178) between the condensers and the collecting vessels, **b**) valves for releasing

170 pressure (v-202, v-179, and v-183) into the ventilation line in case of non-atmospheric

171 experiments and **c**) valves (v-203, v-191, and v-190) beneath the collecting vessel to sample

172 products. The temperature of the first, second, and third condensers are set at $-10 \text{ }^\circ\text{C}$. The

173 outlet of the third condenser is connected to a three-way valve which directs the off-gasses

174 either to a back-pressure regulator ([Figure 1, 8](#)) (EQUILIBAR, U.S.A) or a vacuum pump ([Figure](#)

175 [1, 9](#)) (KNF, SC 920G, Germany). Depending on the desired pressure level, either the vacuum

176 pump or the back-pressure regulator is used to reach vacuum or atmospheric pressure in the
177 reactive, and condensation sections. Subsequently, the outflow of the vacuum pump or the
178 back-pressure regulator (Figure 1, PR-010) is forwarded to the knock-out drum to capture any
179 remaining pyrolyzate. A sampling port was used to collect the gas sample using a Tedlar bag.
180 A drum-type gas flowmeter (Figure 1, 10) (Ritter, TG3/1-1bar, Germany) is employed to
181 measure the volume of the gaseous products before venting them. All pipes, and fittings are
182 made of stainless steel 316 (Swagelok, U.S.A.).

183 2.2.2 Operational procedure and sampling

184 At the beginning of an experiment, the apparatus was cleaned and tested for leak-tightness.
185 The apparatus was flushed with N₂ flow for a sufficiently long time to ensure an inert
186 atmosphere in the unit. The temperature of the condenser's cooler was set at a level of -10
187 °C. The temperature of the extruder's heaters was set at ≤ 300 °C to melt and feed PS without
188 degradation of the PS in the extruder and the four heaters of the extruder were set from the
189 inlet side to the outlet side at 150, 250, 300, and 300 °C, respectively. This temperature profile
190 of the extruder was applied for all experiments. The heat tracing of the reactors' inlet and
191 outlet lines was maintained at 300°C.

192 Next, the pre-cleaned reactor was pre-heated to the desired temperature level. As soon as
193 the temperature at the head of the reactor exceeded the melting point (PS_{m.p.}=240 °C) of the
194 plastics feedstock by 50 °C, i.e. at 290 °C, the plastic feeding was started by turning on the
195 extruder's motor gradually. The feeding rate of the PS feedstock was calibrated once by
196 measuring the required time to feed 1 kg of plastics at three rotational speed settings of the
197 extruder's screw at 10, 20, and 30 revolutions per minute (R.P.M). The R.P.M. was increased
198 gradually until reaching the feeding rate of 1 kg·h⁻¹. The rotational speed of the reactor stirrer

199 was set at 100 R.P.M. to mix the melted plastics in the reactor and to enhance the heat transfer
200 (Figure S1). In each experiment, 4 kg of end-of-life PS feedstock were pyrolyzed. The liquid
201 pyrolyzate was cooled down to -10 °C in the condensation section that comprises 3 “tube in
202 tube” heat exchangers and was collected with an interval of 30 min. The sampling time interval
203 was assumed to be long enough to minimize pressure fluctuations in the reactor. The liquid
204 pyrolyzate samples of each experiment were accumulated in one aluminum bottle and at the
205 end of the experiment, the overall collected liquid pyrolyzate was weighed on a scale
206 (Sartorius ENTRIS8201-1S, Germany). The weight of the gas products was calculated by
207 multiplying the measured volume by the density of the gaseous products. On the one hand,
208 the gaseous product volume was measured using the wet Ritter flowmeter during the entire
209 experiment. On the other hand, the detailed composition of the gaseous products was
210 analyzed using 1D GC-FID/TCD (so-called refinery gas analyzer (RGA)) that has already been
211 explained in detail (Djokic et al., 2017). The analysis was triplicated and the density of the
212 gaseous samples was determined based on the compounds’ wt.% in the gas sample and the
213 pure components’ density. The char yield was calculated by difference. All product yields in
214 the present work were calculated based on the mass of PS fed into the reactor.
215 The mean residence time ($t_{res.}$) is calculated using Equation 1 (Murata et al., 2004).

$$Residence\ time\ [min] = \frac{\text{reactor content [g]}}{\text{feeding rate } [\frac{g}{min}]} \quad \text{Equation 1}$$

216 For each experiment, the reactor content was calculated at each time of sampling by the
217 difference between the fed PS into the reactor and the collected pyrolyzate products,
218 subsequently the $t_{res.}$ was calculated (Equation 1). At the end of each experiment, the average
219 of $t_{res.}$ was calculated.

220 After feeding 4 kg of PS, the feeding was stopped; the pyrolyzate was sampled and the
221 temperature in the reactor was increased up to 600 °C for 30 min to remove all hydrocarbon
222 residues in the reactor before shutting off the coolers and heating elements. The sampled
223 pyrolyzate during the elevated temperature (600 °C) was used only for the mass balance
224 calculation. The rest of the collected samples of each experiment were stored in one
225 aluminum bottle from which the GC × GC sample was taken, and a 4-tert-Butylcatechol (TBC)
226 inhibitor was added to prevent the styrene auto-polymerization.

227 2.3 Experimental conditions

228 Seven PS pyrolysis experiments were design and performed (see [supplementary information](#)).
229 At pressure level of 0.02 bara, three experiments, namely Exp. 1, Exp. 2, and Exp. 3, were
230 performed, at three different temperatures of 450 °C, 550 °C, and 600 °C, respectively. At a
231 pressure of 1.0 bara, the three experiments Exp. 5, Exp. 6, and Exp. 7 were performed at
232 temperatures of 450 °C, 550 °C, and 600 °C, respectively. The 4th experiment (Exp. 4) was
233 operated at a temperature of 500 °C, and a pressure of 0.5 bara. The latter experiment
234 constitutes the midpoint of the operating conditions of Exp. 1, Exp. 2, Exp. 5, and Exp. 6. The
235 4th Exp. was performed to assess the linearity of the yield compositions response with the
236 operating pressure, and temperature. Furthermore, collecting the samples under steady-state
237 conditions was checked, and assured by collecting, and analyzing three samples after 30, 60,
238 and 90 min. The feeding rate was kept constant for all experiments at 1 kg·h⁻¹ and 4 kg of
239 polystyrene was fed for each condition.

240 2.4 Analytical methods

241 For the qualitative analysis, a GC × GC-TOF-MS was used to identify unknown compounds in
242 the liquid pyrolyzate. GC × GC-FID was used to quantify the liquid pyrolyzate composition.

243 Moreover, the composition of the non-condensed off-gasses was analyzed using a 1D GC-
244 FID/TCD (RGA), which is explained in detail in previous work ([Djokic et al., 2017](#)).

245 2.4.1 Sample preparation

246 Two analytes of each pyrolyzate sample were prepared for analysis using GC × GC-FID and GC
247 × GC-ToF-MS. The first analyte was prepared to quantify the highest concentrated
248 compounds, namely, toluene, alpha-methylstyrene, and styrene. The first analytes were
249 prepared by adding 40 wt.% of 3-chlorothiophene to measure the compounds of high
250 concentration. External calibration was performed to determine the response factor of the
251 major compounds such as benzene, toluene, and styrene using high purity chemicals (Sigma-
252 Aldrich, Belgium, purity 99.8%). The second analytes were prepared by adding 5 wt.% of 3-
253 chlorothiophene internal standard, for quantification of the compounds present at lower
254 concentrations. In order to decrease the viscosity and inhomogeneity of the samples, carbon
255 disulfide (CS₂) was used to dilute the sample (1:1 volumetric ratio).

256 2.4.2 GC × GC-FID / TOF-MS setup

257 Two Thermo Scientific TRACE GC × GC instruments (Interscience, Belgium) equipped with an
258 FID detector and a TOF-MS detector (Interscience, Belgium) were used to analyze all samples.
259 The GC × GC was equipped with a dual-stage cryogenic (liquid CO₂) modulator, and a
260 programmable temperature vaporization (PTV) injector (Interscience, Belgium). A non-polar
261 RTX[®]-1 PONA column (Restek, 50 m × 0.25 mm × 0.5 μm) was used as the first dimension
262 column, while a polar BPX-50 column (SGE Analytical Science, 2 m × 0.15 mm × 0.15 μm) was
263 used as the second-dimension column. The PTV temperature was increased from 40 °C up to
264 the maximum temperature of 370 °C with a rate of 15 °C·s⁻¹. The initial GC oven temperature
265 was 40 °C, and it was increased up to 370 °C at a rate of 3 °C·min⁻¹ and held for 600 sec with

266 a set modulation time of 6 sec. The helium carrier gas flows were set to 2.1 mL·min⁻¹ and 2.3
267 mL·min⁻¹ for FID and TOF-MS analysis, respectively (Beens et al., 2005).

268 **Data acquisition and quantification:** Xcalibur™ Software (Thermo Scientific, U.S.A) was used
269 for the acquisition and processing of GC × GC-ToF-MS data. For the GC × GC-FID data, Thermo
270 Scientific's Chrom-Card data system was used. The raw GC × GC-FID data was exported as .cdf
271 file and subsequently processed by GC Image software (Zoex Corporation, U.S.A). The
272 obtained peaks were identified using ToF-MS spectra in comparison with the spectra available
273 in the MS libraries. The blob names and peak volumes were exported as .csv files which were
274 subsequently post-processed. In previous work, the quantification procedure was described
275 in detail by Dijkmans et al. (Dijkmans et al., 2015).

276 The weight fraction wt.%_{*i*} of each compound (*i*) was assigned based on the known weight
277 fraction wt.%_{IS} of internal standard (3-chlorothiophene) using the following equation:

$$wt.\%_i = \frac{f_i \cdot V_i}{f_{IS} \cdot V_{IS}} \cdot wt.\%_{IS} \quad \text{Equation 2}$$

278 *f_i* is the relative response factor of compound *i*, *V_i* is the peak volume of compound *i*, *f_{IS}* is the
279 relative response factor of the internal standard, and *V_{IS}* is the peak volume of the internal
280 standard. The relative response factor was calculated with respect to methane as follows:

$$f_i = \frac{M_i [kg/mol]}{M_{CH_4} [kg/mol] \cdot N_{C,i}} \quad \text{Equation 3}$$

281 where *M_i* is the molar mass of compound *i*, *M_{CH₄}* is the molar mass of methane and *N_{C,*i*}* is the
282 carbon number of compound *i* (Beens et al., 1998).

283 The response factors *f_i* of some compounds (benzene, toluene, and styrene) were determined
284 experimentally. The experimentally measured relative response factors of the mono-aromatic

285 compounds were 90.5% of the calculated response factor based on the effective carbon
286 number. The relative response factors of the remaining compounds in the samples were
287 calculated using the effective carbon number approach used by Djokic et al. (Djokic et al.,
288 2013).

289 2.5 Data visualization and interpolation

290 The Akima mathematical method of interpolation was used for the set of experimental data
291 points (Akima, 1970, 1974, 1978). The z value is interpolated with a bivariate fifth-degree
292 polynomial Equation 4.

$$z(x,y) = \sum_{i=0}^5 \sum_{j=0}^{5-i} a_{ij}x^i y^j \quad \text{Equation 4}$$

293 The interpolation was performed to visualize the effect of the operating parameters pressure
294 (x), and temperature (y) over the pyrolyzate product compounds (z). For each product type,
295 the experimental pyrolyzate results at all operating pressure and temperature levels are used
296 to interpolate points of a mesh of 15 × 15 size. This mathematical model has been used to
297 further interpret the data.

298 3 Results and discussion

299 3.1 Mass balance

300 The yields of the main product fractions obtained using the CSTR continuous pyrolysis reactor
301 are shown in Figure 2. The major product fraction was liquid pyrolyzate with a weight
302 percentage varying between 88.5 and 94.1 wt.%. Under vacuum (0.02 bara) and at
303 temperatures of 450, 550 and 600 °C (Exp. 1, Exp. 2 and Exp. 3), the liquid pyrolyzate yields
304 improved insignificantly at 0.02 bara and it was 91.6, 94.5, and 88.9 wt.% compared to 90.4,

305 94.1 and 88.5 wt.% at 1.0 bara at the corresponding temperatures (Exp. 5, Exp. 6 and Exp. 7).
306 Our experimental results show a trend of increasing liquid pyrolyzate yields with an increased
307 temperature up to 550 °C. An increased temperature above 550 °C led to lower liquid
308 pyrolyzate due to favored secondary reactions at the higher temperature. Similar behavior
309 was observed for PS pyrolysis experiments in a conical spouted bed ([Artetxe et al., 2015](#);
310 [Karaduman et al., 2001](#)) and this behavior was found as well for PS pyrolysis experiments using
311 a fluidized bed reactor at a temperature range from 505 to 782 °C ([Park et al., 2020](#)). This
312 trend can be attributed to reactions of contrary effects that influence the pyrolysis liquid yield
313 recovery: on the one hand, the end-chain β -scission and polymer volatilization enhance the
314 liquid yield recovery with increasing temperature ([Murata et al., 2004](#)). On the other hand,
315 the secondary reactions decrease the pyrolysis liquid yield and stimulate gaseous and char
316 formation with increasing temperature ([Artetxe et al., 2015](#); [Liu et al., 2000](#)).

317 In the studied pressure range (0.02-1.0 bara), the lower pressure (0.02 bara) has an
318 insignificant effect on the yield of the liquid pyrolyzate compared to 1.0 bara at all
319 corresponding temperature levels. In contrast, the vacuum pressure affects significantly the
320 selectivity of the compounds in the liquid yield (see [section 3.3](#)).

321 At the atmospheric pressure level, the total yield of off-gasses is increased from 0.8 at 450 °C
322 (Exp. 5) to 1.5 wt.% at 550 °C (Exp. 6) and finally up to 3.2 wt.% at 600 °C (Exp. 7), the latter as
323 a result of the enforced secondary over-cracking reactions. A similar trend was found at a
324 pressure level of 0.02 bara compared to atmospheric pressure, but with a lower yield at the
325 corresponding temperature levels, which is attributed to the shorter $t_{res.}$ at lower operating
326 pressure.

327 At the pressure level of 0.02 bara, in contrast with liquid pyrolyzate, the char formation
328 decreased from 6.7 wt.% at 450 °C to 3.2 wt.% at 550 °C, then again increased to 8.8 wt.% at
329 550 °C. Likewise, at the atmospheric pressure level, the char formation decreased from 8.9
330 wt.% at 450 °C to 4.4 wt.% at 550 °C, then again increased to 8.3 wt.% at 550 °C. A similar
331 trend was observed by Park et al. and the experimental results of PS pyrolysis in a circulating
332 fluidized bed show a decrease of the char formation before increasing again in a temperature
333 range from 515 to 782 °C (Park et al., 2020). The char formation trend behaves oppositely
334 relative to the pyrolysis liquid formation trend. Enhanced end-chain β -scissions and polymer
335 volatilization reactions at increased temperature are responsible for the increased liquid
336 pyrolysis products and decrease the char formation. On the other hand, the secondary
337 cracking reactions become dominant at temperatures higher than 550 °C, which leads to
338 increased gas yield and char formation and thus a lower liquid yield. The lower pressure
339 minimizes the $t_{res.}$, which mitigates the secondary cracking reactions; as a result, the liquid
340 yield increases.

341 The operating pressure in the studied range had a lower influence on the product yields
342 compared to the operating temperature; In all cases, at operating levels of 0.02 and 1.0 bara,
343 the difference between the product yields was ± 1.3 wt.% at the corresponding temperature
344 level. As an example, at 550 °C, liquid yields at 0.02 and 1.0 bara were 94.5 and 94.1 wt.%,
345 respectively. At the mid-point (Exp. 4, 500 °C and 0.5 bara), the pyrolysis liquid product yield
346 matched 99% of the product yield averages of Exp. 1 (450 °C, 1.0 bara), 2 (550 °C, 0.02 bara),
347 Exp. 5 (450 °C, 1.0 bara) and Exp. 6 (550 °C, 1.0 bara).

348 3.2 The residence time

349 The residence time ($t_{res.}$) influences the rate of secondary reactions and the shorter $t_{res.}$ leads
350 to a lower rate of secondary reactions. At both pressure levels, viz. 0.02 and 1.0 bara, the $t_{res.}$
351 decreases with increased temperature. This observation can be explained by enhanced
352 devolatilization and β -scission reactions (Artetxe et al., 2015; Madorsky, 1952; Murata et al.,
353 2002). At 0.02 bara, the $t_{res.}$ decreased from 39.0, 16.0 to 15.6 min at 450, 550 and 600 °C,
354 respectively. Compared to a reactor pressure of 0.02 bara, the trend was found as well at the
355 1.0 bara level, but higher at the corresponding temperature and decreased from 47.8, 41.6,
356 and 18.6 min at 450, 550, and 600 °C, respectively. The operating temperature has a higher
357 influence compared to the operating pressure on the $t_{res.}$ (see supplementary information).
358 Murata et al. found a similar trend of decreasing $t_{res.}$ with the decreased pressure, however,
359 the authors reported a more pronounced effect in their study due to the higher pressure range
360 from 1.0 bara to 8.0 bara; Murata et al. stated that the degradation and volatilization reactions
361 were a function of temperature, pressure and $t_{res.}$ (Murata et al., 2004). To calculate the $t_{res.}$
362 (Equation 1), the reactor content was estimated by weight difference between the fed PS into
363 the reactor and the sampled pyrolyzate; this procedure can be enhanced further by the
364 addition of a sensor to measure the reactor content in future experimental work.

365 3.3 Composition of pyrolyzate

366 **Comprehensive GC × GC-FID analysis:** in all cases, over 95 wt.% of the collected liquid fractions
367 were quantified and the dominant compound was styrene. The carbon number of the
368 detected compounds ranged from C₃ to C₂₄. The GC × GC-FID analyses results showed on
369 average major amounts of mono-aromatics (62 wt.%), di-aromatics (13 wt.%), and tri-
370 aromatics (7 wt.%) (Table 1), as well as small amounts of tetra-aromatics and aliphatics.

371 [Figure 3](#) shows the comprehensive GC × GC-FID chromatogram of the end-of-life PS pyrolyzate
372 (Exp. 1). The chromatogram shows five aromatic groups (mono-aromatics, di-aromatics, tri-
373 aromatics, tetra-aromatics, and penta-aromatics) and the aliphatic group. The key compounds
374 of end-of-life PS pyrolyzate are indicated in the chromatogram ([Figure 3](#)). The major
375 compounds of the mono-aromatic groups were **(c)** Toluene **(e)** Ethylbenzene **(f)** Styrene **(j)**
376 Alpha-methylstyrene. The dominant compounds of the poly-aromatic groups were **(q)**
377 Bibenzyl **(r)** 1,2-Diphenylpropane **(t)** 1,3-Diphenylpropane **(u)** 2,4-Diphenyl-1-butene **(v)** 1,3-
378 Diphenylpropene **(w)** 2,4-Diphenyl-1-pentene **(x)** 2,4,6-triphenyl-1-hexene **(y)** 1,2,4-
379 Triphenylbenzene. The polypropylene contamination in the PS feedstock is detected by
380 identifying the 2, 4, 6, 8-tetramethyl-1-undecene in the pyrolyzate—one of the major
381 fingerprint compounds from polypropylene pyrolysis ([Soják et al.](#)). Additionally, small
382 amounts of compounds with oxygen and/or nitrogen atoms were detected, such as Benzene-
383 acetamide, which is attributed to the impurity of the end-of-life PS feedstock.

384 3.3.1 Composition of pyrolyzate at the atmospheric pressure

385 [Figure 4](#) shows the yield of the four major groups found in the liquid pyrolyzate: i) styrene, ii)
386 other mono-aromatics, iii) poly-aromatics (PA) and iv) aliphatic compounds. In all investigated
387 process conditions, the major compound of the pyrolyzate was styrene. The yield of styrene
388 increased from 35.7 wt.% at 450 °C to 41.2 wt.% at 550 °C and reached 43.2 wt.% at 600 °C.
389 Liu et al. studied the effect of temperature on polystyrene pyrolysis yield from 450 °C to 700
390 °C and found that the styrene yield increased with the increased temperature to reach a
391 certain temperature point (600 °C) at which the styrene concentration was maximum (78.8
392 wt.%), then decreased to 60 wt.% at 700 °C ([Liu et al., 2000](#)). This trend can be attributed to
393 reactions of contrary effects that influence the styrene production: on the one hand, the

394 styrene concentration increases because of the enhanced depolymerization rate and end-
395 chain β -scission reactions with the increased operating temperature (Artetxe et al., 2015;
396 Madorsky, 1952; Murata et al., 2002). On the other hand, the secondary reactions decrease
397 the styrene concentration and stimulate gases and char formation with increasing
398 temperature (Artetxe et al., 2015; Liu et al., 2000). In this work, the styrene concentration
399 increased with the increased temperature until reaching the maximum operating temperature
400 of 600 °C. The temperature point of maximum styrene is assumed to lie beyond the maximum
401 studied temperature level (600 °C). The temperature level was restricted because it became
402 difficult to sustain a stable operating temperature at higher levels.

403 At atmospheric pressure, the yield of other mono-aromatics (excluding styrene) decreased
404 from 25.1 wt.%, 25.0 wt.% to 22.9 wt.% with increased temperature from 450 °C, 550 °C to
405 600 °C, respectively, in favor of increased styrene yields (Figure 4).

406 On the other hand, the yield of benzene, toluene, ethylbenzene, and xylenes (BTEX) decreased
407 from 17.1 wt.% at 450 °C (Exp. 5) to 15.6 wt.% at 550 °C and 600 °C (Exp. 6, Exp. 7). Similarly,
408 the alpha-methylstyrene decreased from 6.1 wt.% at 450 °C (Exp. 5), 6.0 wt.% at 550 °C (Exp.
409 6) to 4.4 wt.% at 600 °C (Exp. 7) (Figure 4 and Figure 5). The decrement of these mono-
410 aromatics yields is attributed to the augmented secondary reactions. In contrast, it was
411 reported that in a fluidized bed reactor, the recovery of BTEX increased from 1.9 to 6.9 wt.%
412 with the increased temperature from 515 to 628 °C (Park et al., 2020). As a result, the process
413 and apparatus used in the present work have the superiority of producing BTEX at the
414 operating temperature of less than 600 °C. Only at higher operating temperatures of 698 and
415 782 °C, Park et al. obtained a higher BTEX yield (26.3 wt.%) compared to the present work
416 (Park et al., 2020).

417 **Poly-aromatics:** The yield of poly-aromatics decreased from 21.6 wt.% at 450 °C (Exp. 5), 21.0
418 wt.% at 550 °C (Exp. 6) to 15.8 wt.% at 600 °C (Exp. 7) in favor of an increased styrene yield; a
419 similar trend of decreased poly-aromatics with increased temperature was found by Artetxe
420 et al. (Artetxe et al., 2015). Two primary reactions may affect the formation of higher poly-
421 aromatic products: secondary cracking reactions and recombination reactions. The
422 recombination reactions of large radicals increase the yield of poly-aromatic products,
423 whereas the secondary (over-)cracking reactions decrease the yield of higher poly-aromatics
424 products. It should be noted that the elevated temperature increases the secondary (over-
425)cracking reactions and the termination reactions at different rates. At a certain temperature
426 level (optimum temperature), the rate of the secondary cracking reactions exceeds the rate
427 of the recombination reactions. Subsequently, the formation of poly-aromatic products
428 decreases, as shown in Figure 4. Furthermore, the formation of mono-aromatics such as
429 benzene, toluene, and styrene increased with the increased temperature (Figure 5) which
430 agrees with the findings of Liu et al. (Liu et al., 2000). Finally, the yield of the aliphatic
431 compound was 5.2 ± 1.9 wt.%. The GC \times GC analysis confirms that the aliphatic compounds
432 were highly branched and it is likely that polypropylene contamination is the major source of
433 the aliphatic compounds.

434 3.3.2 Composition of pyrolyzate at reduced pressures (0.02 and 0.5 bara)

435 The reduced pressure affected the PS pyrolysis yield and enhanced the styrene recovery (see
436 Figure 4 and Table 1). Furthermore, the styrene had the highest concentration at 0.02 bara
437 pressure level compared with the corresponding temperature at the 1.0 bara pressure level.
438 Chauhan et al. pyrolyzed polystyrene at reduced pressure through microwave-assisted
439 pyrolysis and found a similar trend of increased styrene yield at vacuum pressure (Chauhan et

440 [al., 2008](#)). Vacuum minimizes the secondary cracking reactions effect and thus maximizes the
441 primary product styrene due to a shorter $t_{res.}$ in the reactor (see, section 3.2); which is in line
442 with Bartoli et al.'s PS pyrolysis experiments findings ([Bartoli et al., 2015](#)). The styrene yield of
443 the mid-point Exp. 4 (500 °C and 0.5 bara) is 41.7 wt.% and it is 2.8 wt.% lower than the average
444 styrene yield of Exp.s 1, 2, 5 and 6 (44.5 wt.%). As such, it appears the effect of pressure on
445 the styrene yield is not completely linear, even though the effect of pressure on the total liquid
446 yield was fairly linear. Moreover, the styrene yield increases with the increased temperature
447 (Figure 4). Likewise, Yang and Shibasaki conducted polystyrene pyrolysis experiments using
448 Py-GC and found an increment of styrene yield with increased temperature ([Yang and
449 Shibasaki, 1998](#)). But, beyond the temperature point (550 °C) at which the styrene
450 concentration is maximum (55.9 wt.%), the styrene yield dropped due to enhanced secondary
451 (over-)cracking reactions of intermediate products ([Liu et al., 2000](#)). Consequently, the
452 secondary reaction products such as gaseous products (C₁-C₄) increased with the increased
453 temperature ([Artetxe et al., 2015](#)). Furthermore, the improvement of styrene concentration
454 at vacuum pressure (0.02 bara) was 14.7 wt.% at (Run 2) 0.02 bara and 550 °C compared to
455 Run 6) 1.0 bara and 550 °C. Besides improving the styrene recovery, the vacuum pressure
456 shifted the temperature point at which the styrene concentration is maximum to the lower
457 level, namely 550 °C. This is could be attributed to the combined effects of pressure and
458 temperature, which could lead to different response rates of the reactions of contrary effects:
459 1. secondary reactions and 2. depolymerization and β -scission reactions.

460 **Other Mono-aromatics:** At vacuum conditions, the yield of other mono-aromatics (excluding
461 styrene) accounts for 9.6 ± 1.3 wt.%, which was lower than the yield at atmospheric pressure
462 at all corresponding temperature levels (450, 550 and 600 °C) ([Figure 4](#)). The decreased other

463 mono-aromatics compounds under vacuum is attributed to the shorter t_{res} , which accelerates
464 the removal of primary products and reduces secondary reactions. The yield of other mono-
465 aromatics at the mid-point Exp. 4 is 18.9 wt.% and 1.6 wt.% higher than in experiments 1, 2,
466 6, and 7 (17.3 wt.%).

467 The yield of BTEX increased slightly from 4.3 wt.% at 450 °C (Exp. 5) to peak at 5.6 wt.% at 550
468 °C (Exp. 6) and then decreased to 4.9 wt.% at 600 °C (Exp. 7) (Figure 5); which is in line with
469 Artetxe et al.'s findings on increased single-ring aromatic products (such as toluene,
470 ethylbenzene, and α -methylstyrene) with an increased temperature from 450 to 550 °C, while
471 a further increase above 550 °C led to a decrease in their yield (Artetxe et al., 2015). Similarly,
472 the yield of α -methylstyrene changed slightly between 3.2 and 3.4 wt.% (Figure 5). Under
473 vacuum, the BTEX and α -methylstyrene yields were reduced compared to atmospheric
474 pressure at the corresponding temperature levels due to the decreased secondary reaction
475 under shorter t_{res} .

476 **Poly-aromatics:** On the one hand, the dimers such as 2,4-Diphenyl-1-butene are formed via
477 1,3-hydrogen transfer reactions followed by mid-chain β -scission reactions (Huang et al.,
478 2020; Levine and Broadbelt, 2008). The dimer compound (2,4-Diphenyl-1-butene) was slightly
479 affected and decreased from 3.8 to 3.7 wt.% with the increased temperature from 450 to 600
480 °C. A similar minor effect of temperature on the dimer formation was found by Artetxe et al.
481 (Artetxe et al., 2015). On the other hand, styrene trimer (2,4,6-triphenyl-1-hexene) is formed
482 via 1,5-hydrogen transfer reactions followed by mid-chain β -scission reactions (Levine and
483 Broadbelt, 2008). And the increased temperature from 450 to 600 °C affected drastically the
484 formation of 2,4,6-triphenyl-1-hexene, which decreased from 12.8 to 0.1 wt.% due to de-
485 polymerization steps and secondary reactions (Artetxe et al., 2015).

486 The yield of poly-aromatics (higher than di-aromatics and tri-aromatics) decreased from 29.5
487 wt.% at 450 °C (Exp. 5) to 22.3 wt.% at 550 °C (Exp. 6) and further to 15.2 wt.% at 600 °C (Exp.
488 7). The yield of poly-aromatics from experiments under vacuum was higher compared to the
489 atmospheric pressure experiments at the corresponding temperature. The vacuum shortens
490 the t_{res} , which minimizes cracking of poly-aromatics compared with atmospheric pressure
491 (Figure 4). The mitigated secondary cracking reactions under vacuum resulted in a smaller
492 amount of other mono-aromatics. Finally, the yield of aliphatic compounds was 5.6 ± 1.6 wt.%
493 which is in the same range as the yield of aliphatic products at atmospheric pressure (Figure
494 4).

495 Table 1 summarizes the product yields from pyrolysis of end-of-life PS at the different pressure
496 and temperature levels. In all cases, the gas yield counts less than 4 wt.% and the major
497 products are liquid with a minimum yield of 88.5 wt.%. The impurity of the end-of-life
498 polystyrene feedstock was apparent through detecting heteroatomic compounds in the
499 pyrolyzate such as benzene-acetamide; in contrast, these compounds were not detected in
500 previous works that used virgin or ultra-pure end-of-life PS (Achilias et al., 2007; Artetxe et al.,
501 2015; Park et al., 2020).

502 In the aim of comparing the effect of feedstock purity on the composition of the pyrolyzate, a
503 virgin PS was processed at 600 °C and 0.02 bara, i.e. under the same operating conditions of
504 Exp. no. 3. The results can be seen in Figure 6.

505 Processing virgin PS resulted in a styrene yield of 67.1 wt.%, compared to only 54.5 wt.% of
506 styrene in the end-of-life PS pyrolysis case (Figure 6). This is attributed to the differences in
507 feedstock purity, where quantitatively, the contamination of the end-of-life PS feedstock

508 decreases the net PS in the processed end-of-life PS sample. Furthermore, metallic
509 contaminations such as Mg, Al, Na, and Ca (Roosen et al., 2020b) may affect the pyrolysis
510 process qualitatively. The metallic contaminations play a role as pyrolysis catalyst, which
511 explains the lower liquid pyrolysis yield as found by Iftikhar et al. (Iftikhar et al., 2019). Note
512 that the char formation of end-of-life PS pyrolysis increased by 1.8 wt.% compared to virgin
513 PS pyrolysis case; this behavior can be attributed as well to the metallic contamination
514 contents of the end-of-life PS which favors higher char formation compared to using virgin PS
515 without metallic contaminations (Kabir and Hameed, 2017; Lin et al., 2018). Also, in the end-
516 of-life PS pyrolysis, the mono-aromatics (excluding styrene) yield increased by 4.7 wt.%
517 compared to the virgin PS pyrolysis case because of the metallic contamination's presence
518 and impurities such as PET, PA, and PUR which may decompose into aromatics. The aliphatic
519 compounds yield was 7.1 wt.% higher with end-of-life PS because of the impurity of the
520 feedstock with other plastics types such as polyolefins (Roosen et al., 2020b), whereas no
521 aliphatic compounds were detected in the virgin PS pyrolyzate. The gas products (C₁-C₄)
522 increased slightly from 1.86 wt.% with virgin PS pyrolysis to 2.3 wt.% with end-of-life PS
523 pyrolysis as the feedstock.

524 3.4 Optimization of pyrolyzate yield and composition

525 A secondary objective of this study was set to optimize the process conditions depending on
526 the desired product (e.g. styrene versus benzene or dimer yield). Figure 7 (a) shows that the
527 pressure has a limited effect on the liquid pyrolyzate yield, while the temperature increases
528 the liquid pyrolyzate yield up to 94.5 wt.% at 550 °C and 0.02 bara. At higher temperature
529 levels, the liquid pyrolyzate yield decreases again due to secondary and over-cracking
530 reactions that favor gas (C₁-C₄) and char formation (Figure 7 (a)). Note that the contour plot

531 of liquid pyrolyzate can be used as a guideline to calculate the cost of pyrolyzate production
532 depending on the operating cost at varied pressure and temperature levels. Moreover, the
533 optimum operating conditions of the pyrolyzate liquid yield could differ from the optimum
534 operating conditions for styrene. [Figure 7 \(b\)](#) shows that the styrene yield reaches maxima
535 between 550 and 575 °C at 0.02 bara with a styrene yield of 56 wt.%. Higher pressure (1.0
536 bara) and lower temperatures (450 °C) decrease the styrene yield substantially, to less than
537 36 wt.% ([Figure 7 \(b\)](#)). The pressure has no actual effect on the liquid yield but a large effect
538 on the styrene concentration ([Figure 7 \(a, b\)](#)). The yield of benzene, toluene, ethylbenzene,
539 and xylenes (BTEX) vary between 8.4 and 27.6 wt.% ([Figure 7 \(c\)](#)). In the studied pressure and
540 temperature ranges, the pressure has a more pronounced effect compared to the
541 temperature on the yield of BTEX.

542 [Figure 7 \(d\)](#) shows the ratio of styrene to BTEX (St./BTEX) at different pressure and
543 temperature levels. This parameter can be used to tune the process based on the market
544 requirements and/or the installation and operating costs of the distillation equipment. The
545 lowest ratio of St./BTEX (1.6) was obtained at a 1.0 bara pressure level and temperature level
546 of 500 °C. The St./BTEX ratio increased up to 5.6 with increased temperature in the range of
547 560 °C and 600 °C and lower pressure (0.02 bara).

548 The yield of trimers and dimers can be optimized depending on the industrial and market
549 conditions as well. The dimer and trimer yield increased with increased pressure and
550 temperature ([Figure 7 \(e\)](#)) and varied in the range of 5 to 17 wt.%. Both increased pressure
551 and temperature reduced the yield of dimers and trimers due to the increased $t_{res.}$ with
552 increased pressure and augmented secondary cracking reactions. Finally, the yield of higher
553 poly-aromatics excluding di-aromatics and tri-aromatics is shown in ([Figure 7 \(f\)](#)). The minimal

554 yield of this fraction is at a higher temperature level of 600 °C at both pressure levels from
555 0.02 to 1.0 bara, which is attributed to the shorter t_{res} . and the dominance of the secondary
556 cracking reaction. The obtained results can be used as an input for a comprehensive techno-
557 economical assessment. Moreover, the process of PS pyrolysis can be further optimized based
558 on the target components and feedstock purity. As mentioned earlier, an increase in
559 temperature and decrease in pressure enhances the devolatilization and de-propagation
560 reactions.

561 4 Conclusion

562 The continuous pyrolysis process of end-of-life PS was studied in a vacuum and atmospheric
563 pressure in a continuous stirred tank reactor (CSTR). A maximum yield of benzene, toluene,
564 ethylbenzene, and xylene BTEX (17.1 wt.%) was observed at 450 °C and 1.0 bara, which
565 corresponds to a lower temperature for a similar yield of PS pyrolysis in a fluidized bed reactor.
566 At the pressure level of 0.02 bara, the maximum yield of styrene (55.9 wt.%) was achieved at
567 550 °C which was considered to be the temperature point of maximum yield; beyond of this
568 optimal point, the styrene yield decreased. Compared to the atmospheric pressure level, the
569 vacuum pressure decreased the temperature point from 600 °C to 550 °C which is
570 economically favorable due to lower energy consumption to pyrolyze the PS.

571 The operating pressure and temperature affected the product distribution in the studied
572 ranges of 450-600 °C and 0.02-1.0 bara, but in these ranges, the temperature had a more
573 pronounced effect. The GC × GC-FID and GC × GC-TOF/MS analyses led to a detailed
574 characterization of the polystyrene pyrolyzate. Interestingly, feedstock impurities and
575 contaminations could be qualified by detecting unique pyrolysis product compounds such as

576 2, 4, 6, 8-tetramethyl-1-undecene, which is a characteristic compound of polypropylene
577 pyrolysis. The latter was an indication that PP was a contaminant of end-of-life PS stream.
578 These analyses results show the superiority and necessity of using GC × GC for analyzing a
579 complex contaminated end-of-life PS pyrolyzate and to avoid overlapping of compounds'
580 peaks. A higher purity feedstock led to higher styrene yield, with styrene being the most
581 valuable PS pyrolyzate product. On the other hand, further purification of the end-of-life PS
582 would potentially increase the cost of pre-processing and treatments; therefore, a techno-
583 economical assessment is required to maximize the styrene yield with minimal pre-treatment
584 cost. To ensure the quality and purity of the end-of-life plastic feedstock, comprehensive GC
585 × GC analysis or fast screening methods based on pyrolysis GC can be used to analyze the
586 liquid pyrolysis products of these feedstocks. Since the post-processing and distillation of the
587 liquid pyrolyzate result poly-aromatics cuts that contain dimers and trimers, further studies
588 are therefore necessary to clarify the potential of re-pyrolyzing of the poly-aromatics cuts to
589 produce styrene and BTEX. Measuring the heat input of the extruder and the energy
590 consumption of the auxiliary equipment will help in performing a comprehensive techno-
591 economical assessment. In addition, the research on removing end-of-life PS contaminants by
592 pre-processing steps can be planned for future work. Finally, the presented results for
593 pyrolysis of end-of-life PS can be used as a basis for scale-up to an industrial-sized plant.

594 [Acknowledgments](#)

595 This work was supported by Flanders Innovation & Entrepreneurship (VLAIO) and the O&O
596 project “Thermochemical Conversion of End-of-Life Plastics to Chemical Feedstock”;
597 (Nederlandse vertaling: Thermochemische Omzetting van End-of-Life Kunststoffen tot
598 Chemische Feedstocks). The research leading to these results has also received funding from

599 the Fund for Scientific Research Flanders (FWO) and innovation program/ERC grant
600 agreement no.818607 (OPTIMA).

601 [Declaration of competing interest](#)

602 The authors declare no conflict of interest.

603 [CRediT authorship contribution statement](#)

604 **Azd Zayoud**: conceptualization, methodology, investigation, writing – original draft, writing –
605 review & editing. **Hang Dao Thi**: investigation, writing – review & editing. **Marvin Kusenberg**:
606 methodology, investigation, writing – review & editing. **Andreas Eschenbacher**: writing –
607 review & editing. **Uros Kresovic**: resources, review & editing. **Nick Alderweireldt**: resources,
608 review & editing. **Marko Djokic**: methodology, investigation, writing – review & editing. **Kevin**
609 **M. Van Geem**: conceptualization, funding acquisition, project administration, supervision,
610 writing – review & editing.

611 References

- 612 Achilias, D.S., Kanellopoulou, I., Megalokononimos, P., Antonakou, E., Lappas, A.A., 2007. Chemical
613 Recycling of Polystyrene by Pyrolysis: Potential Use of the Liquid Product for the Reproduction of
614 Polymer. *Macromol Mater Eng* 292, 923-934, <https://doi.org/10.1002/mame.200700058>.
- 615 Agilyx, 2020. Driving innovations to make plastics a renewable resource.
- 616 Akima, H., 1970. A New Method of Interpolation and Smooth Curve Fitting Based on Local Procedures.
617 *J Acm* 17, 589-602, <https://doi.org/10.1145/321607.321609>.
- 618 Akima, H., 1974. A method of bivariate interpolation and smooth surface fitting based on local
619 procedures. *Communications of the ACM* 17, 18-20, <https://doi.org/10.1145/360767.360779>.
- 620 Akima, H., 1978. A Method of Bivariate Interpolation and Smooth Surface Fitting for Irregularly
621 Distributed Data Points. *ACM Transactions on Mathematical Software* 4, 148-159,
622 <https://doi.org/10.1145/355780.355786>.
- 623 Ando, H., Inomata, O., Abe, T., Matsuzawa, S., Shimizu, Y., Miyazaki, S., 1974. Pyrolysis of Plastic Wastes
624 (I). *Journal of the Fuel Society of Japan* 53, 646-653, https://doi.org/10.3775/jie.53.7_646.
- 625 Artetxe, M., Lopez, G., Amutio, M., Barbarias, I., Arregi, A., Aguado, R., Bilbao, J., Olazar, M., 2015.
626 Styrene recovery from polystyrene by flash pyrolysis in a conical spouted bed reactor. *Waste Manag*
627 45, 126-133, <https://doi.org/10.1016/j.wasman.2015.05.034>.
- 628 Bartoli, M., Rosi, L., Frediani, M., Undri, A., Frediani, P., 2015. Depolymerization of polystyrene at
629 reduced pressure through a microwave assisted pyrolysis. *Journal of Analytical and Applied Pyrolysis*
630 113, 281-287, <https://doi.org/10.1016/j.jaap.2015.01.026>.
- 631 Beens, J., Boelens, H., Tijssen, R., Blomberg, J., 1998. Quantitative Aspects of Comprehensive Two-
632 Dimensional Gas Chromatography (GC×GC). *Journal of High Resolution Chromatography* 21, 47-54.
- 633 Beens, J., Janssen, H.G., Adahchour, M., Brinkman, U.A., 2005. Flow regime at ambient outlet pressure
634 and its influence in comprehensive two-dimensional gas chromatography. *J Chromatogr A* 1086, 141-
635 150, <https://doi.org/10.1016/j.chroma.2005.05.086>.
- 636 Brooks, A.L., Wang, S., Jambeck, J.R., 2018. The Chinese import ban and its impact on global plastic
637 waste trade. *Sci Adv* 4, eaat0131, <https://doi.org/10.1126/sciadv.aat0131>.
- 638 Chauhan, R.S., Gopinath, S., Razdan, P., Delattre, C., Nirmala, G.S., Natarajan, R., 2008. Thermal
639 decomposition of expanded polystyrene in a pebble bed reactor to get higher liquid fraction yield at
640 low temperatures. *Waste Manag* 28, 2140-2145, <https://doi.org/10.1016/j.wasman.2007.10.001>.
- 641 Dalluge, J., van Rijn, M., Beens, J., Vreuls, R.J., Brinkman, U.A., 2002. Comprehensive two-dimensional
642 gas chromatography with time-of-flight mass spectrometric detection applied to the determination of
643 pesticides in food extracts. *J Chromatogr A* 965, 207-217, [https://doi.org/10.1016/S0021-9673\(01\)01324-3](https://doi.org/10.1016/S0021-9673(01)01324-3).
- 644
- 645 Dijkmans, T., Djokic, M.R., Van Geem, K.M., Marin, G.B., 2015. Comprehensive compositional analysis
646 of sulfur and nitrogen containing compounds in shale oil using GC×GC – FID/SCD/NCD/TOF-MS. *Fuel*
647 140, 398-406, <https://doi.org/10.1016/j.fuel.2014.09.055>.
- 648 Djokic, M., Carstensen, H.-H., Van Geem, K.M., Marin, G.B., 2013. The thermal decomposition of 2,5-
649 dimethylfuran. *Proceedings of the Combustion Institute* 34, 251-258,
650 <https://doi.org/10.1016/j.proci.2012.05.066>.
- 651 Djokic, M.R., Ristic, N.D., Olahova, N., Marin, G.B., Van Geem, K.M., 2017. Quantitative on-line analysis
652 of sulfur compounds in complex hydrocarbon matrices. *J Chromatogr A* 1509, 102-113,
653 <https://doi.org/10.1016/j.chroma.2017.06.006>.
- 654 Doucet, J., 2020. The Endless Regeneration of Plastics Through Microwaves.
- 655 Doucet, J., Chaouki, J., Sobhy, A., 2016. Catalyst for distributed batch microwave pyrolysis, system and
656 process thereof.
- 657 Garside, M., 2019. Global plastic production from 1950 to 2018 (in million metric tons).

658 Huang, J., Li, X., Meng, H., Tong, H., Cai, X., Liu, J., 2020. Studies on pyrolysis mechanisms of syndiotactic
659 polystyrene using DFT method. Chemical Physics Letters 747,
660 <https://doi.org/10.1016/j.cplett.2020.137334>.

661 Iftikhar, H., Zeeshan, M., Iqbal, S., Muneer, B., Razzaq, M., 2019. Co-pyrolysis of sugarcane bagasse and
662 polystyrene with ex-situ catalytic bed of metal oxides/HZSM-5 with focus on liquid yield. Bioresour
663 Technol 289, 121647, <https://doi.org/10.1016/j.biortech.2019.121647>.

664 Kabir, G., Hameed, B.H., 2017. Recent progress on catalytic pyrolysis of lignocellulosic biomass to high-
665 grade bio-oil and bio-chemicals. Renewable and Sustainable Energy Reviews 70, 945-967,
666 <https://doi.org/10.1016/j.rser.2016.12.001>.

667 Kaminsky, W., 2021. Chemical recycling of plastics by fluidized bed pyrolysis. Fuel Communications 8,
668 <https://doi.org/10.1016/j.jfueco.2021.100023>.

669 Kaminsky, W., Franck, J., 1991. Monomer recovery by pyrolysis of poly(methyl methacrylate) (PMMA).
670 Journal of Analytical and Applied Pyrolysis 19, 311-318, [https://doi.org/10.1016/0165-2370\(91\)80052-](https://doi.org/10.1016/0165-2370(91)80052-)
671 [a](https://doi.org/10.1016/0165-2370(91)80052-a).

672 Kaminsky, W., Predel, M., Sadiki, A., 2004. Feedstock recycling of polymers by pyrolysis in a fluidised
673 bed. Polymer Degradation and Stability 85, 1045-1050,
674 <https://doi.org/10.1016/j.polymdegradstab.2003.05.002>.

675 Karaduman, A., Şimşek, E.H., Çiçek, B., Bilgesü, A.Y., 2001. Flash pyrolysis of polystyrene wastes in a
676 free-fall reactor under vacuum. Journal of Analytical and Applied Pyrolysis 60, 179-186,
677 [https://doi.org/10.1016/s0165-2370\(00\)00169-8](https://doi.org/10.1016/s0165-2370(00)00169-8).

678 Kim, Y.S., Hwang, G.C., Bae, S.Y., Yi, S.C., Moon, S.K., Kumazawa, H., 1999. Pyrolysis of polystyrene in a
679 batch-type stirred vessel. Korean J Chem Eng 16, 161-165, <https://doi.org/10.1007/bf02706830>.

680 Levine, S.E., Broadbelt, L.J., 2008. Reaction pathways to dimer in polystyrene pyrolysis: A mechanistic
681 modeling study. Polymer Degradation and Stability 93, 941-951,
682 <https://doi.org/10.1016/j.polymdegradstab.2008.01.029>.

683 Lin, X., Zhang, Z., Zhang, Z., Sun, J., Wang, Q., Pittman, C.U., 2018. Catalytic fast pyrolysis of a wood-
684 plastic composite with metal oxides as catalysts. Waste Manag 79, 38-47,
685 <https://doi.org/10.1016/j.wasman.2018.07.021>.

686 Liu, Y., Qian, J., Wang, J., 2000. Pyrolysis of polystyrene waste in a fluidized-bed reactor to obtain
687 styrene monomer and gasoline fraction. Fuel Processing Technology 63, 45-55,
688 [https://doi.org/10.1016/s0378-3820\(99\)00066-1](https://doi.org/10.1016/s0378-3820(99)00066-1).

689 Madorsky, S., Straus, S., 1948. Pyrolytic fractionation of polystyrene in a high vacuum and mass
690 spectrometer analysis of some of the fractions. Journal of research of the National Bureau of Standards
691 40, 417-425.

692 Madorsky, S.L., 1952. Rates of thermal degradation of polystyrene and polyethylene in a vacuum.
693 Journal of Polymer Science 9, 133-156, <https://doi.org/10.1002/pol.1952.120090203>.

694 Mo, Y., Zhao, L., Wang, Z., Chen, C.L., Tan, G.Y., Wang, J.Y., 2014. Enhanced styrene recovery from
695 waste polystyrene pyrolysis using response surface methodology coupled with Box-Behnken design.
696 Waste Manag 34, 763-769, <https://doi.org/10.1016/j.wasman.2014.01.005>.

697 Murata, K., Hirano, Y., Sakata, Y., Uddin, M.A., 2002. Basic study on a continuous flow reactor for
698 thermal degradation of polymers. Journal of Analytical and Applied Pyrolysis 65, 71-90,
699 [https://doi.org/10.1016/s0165-2370\(01\)00181-4](https://doi.org/10.1016/s0165-2370(01)00181-4).

700 Murata, K., Sato, K., Sakata, Y., 2004. Effect of pressure on thermal degradation of polyethylene.
701 Journal of Analytical and Applied Pyrolysis 71, 569-589, <https://doi.org/10.1016/j.jaap.2003.08.010>.

702 Ogino, H., Nagy, B., 1981. Pyrolysis of Transvaal kerogens. II. An evaluation of vacuum pyrolysis with
703 polyethylene, polystyrene and their mixtures with minerals. Precambrian Research 15, 113-130,
704 [https://doi.org/10.1016/0301-9268\(81\)90025-5](https://doi.org/10.1016/0301-9268(81)90025-5).

705 Park, J.J., Park, K., Kim, J.-S., Maken, S., Song, H., Shin, H., Park, J.-W., Choi, M.-J., 2003. Characterization
706 of Styrene Recovery from the Pyrolysis of Waste Expandable Polystyrene. Energy & Fuels 17, 1576-
707 1582, <https://doi.org/10.1021/ef030102l>.

708 Park, K.-B., Jeong, Y.-S., Guzelciftci, B., Kim, J.-S., 2020. Two-stage pyrolysis of polystyrene: Pyrolysis oil
709 as a source of fuels or benzene, toluene, ethylbenzene, and xylenes. *Appl Energy* 259,
710 <https://doi.org/10.1016/j.apenergy.2019.114240>.

711 Phillips, J.B., Beens, J., 1999. Comprehensive two-dimensional gas chromatography: a hyphenated
712 method with strong coupling between the two dimensions. *Journal of Chromatography A* 856, 331-
713 347, [https://doi.org/10.1016/s0021-9673\(99\)00815-8](https://doi.org/10.1016/s0021-9673(99)00815-8).

714 PlasticsEurope, 2018. *Plastics the facts 2018*.

715 PlasticsEurope, 2019a. *Plastics 2030 PlasticsEurope's Voluntary Commitment to increasing circularity
716 and resource efficiency*.

717 PlasticsEurope, 2019b. *Plastics the facts 2019*.

718 PolystyreneLoop, 2021. *PolyStyreneLoop project: pioneer demonstration plant to treat up to 3300
719 tonnes of polystyrene waste/yr as of 2018*.

720 Pyrolyze.B.V., 2021. 150 kg/h plastic pyrolysis plant.

721 Qureshi, K.M., Kay Lup, A.N., Khan, S., Abnisa, F., Wan Daud, W.M.A., 2018. A technical review on semi-
722 continuous and continuous pyrolysis process of biomass to bio-oil. *Journal of Analytical and Applied
723 Pyrolysis* 131, 52-75, <https://doi.org/10.1016/j.jaap.2018.02.010>.

724 Qureshi, M.S., Oasmaa, A., Lindfors, C., 2019. <Thermolysis of plastic waste_Reactor comparison.pdf>,
725 in: Berruti, F., Dufour, A., Prins, W., Garcia-Pérez, M. (Eds.), *Pyroliq 2019: Pyrolysis and Liquefaction of
726 Biomass and Wastes*, Cork, Ireland

727 Roosen, M., Mys, N., Kusenberg, M., Billen, P., Dumoulin, A., Dewulf, J., Van Geem, K.M., Ragaert, K.,
728 De Meester, S., 2020a. Detailed Analysis of the Composition of Selected Plastic Packaging Waste
729 Products and Its Implications for Mechanical and Thermochemical Recycling. *Environ. Sci. Technol.* 54,
730 13282-13293, <https://doi.org/10.1021/acs.est.0c03371>.

731 Roosen, M., Mys, N., Kusenberg, M., Billen, P., Dumoulin, A., Dewulf, J., Van Geem, K.M., Ragaert, K.,
732 De Meester, S., 2020b. Detailed Analysis of the Composition of Selected Plastic Packaging Waste
733 Products and Its Implications for Mechanical and Thermochemical Recycling. *Environ Sci Technol* 54,
734 13282-13293, <https://doi.org/10.1021/acs.est.0c03371>.

735 Smalley, M., 2019. Agilyx partners to launch European PS chemical recycling plant: Styrenics Circular
736 Solutions, Agilyx aim to develop a commercial-scale chemical recycling plant.
737 www.recyclingtoday.com.

738 Soják, L., Kubinec, R., Jurdáková, H., Hájeková, E., Bajus, M., 2007. High resolution gas
739 chromatographic–mass spectrometric analysis of polyethylene and polypropylene thermal cracking
740 products. *Journal of Analytical and Applied Pyrolysis* 78, 387-399,
741 <https://doi.org/10.1016/j.jaap.2006.09.012>.

742 Toraman, H.E., Dijkmans, T., Djokic, M.R., Van Geem, K.M., Marin, G.B., 2014. Detailed compositional
743 characterization of plastic waste pyrolysis oil by comprehensive two-dimensional gas-chromatography
744 coupled to multiple detectors. *J Chromatogr A* 1359, 237-246,
745 <https://doi.org/10.1016/j.chroma.2014.07.017>.

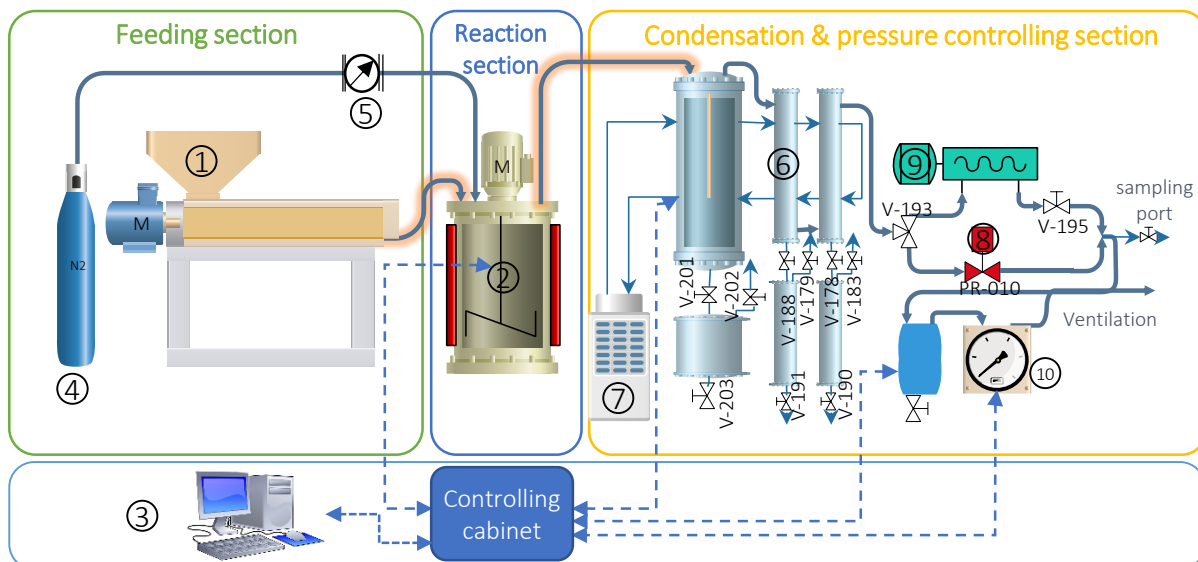
746 Undri, A., Frediani, M., Rosi, L., Frediani, P., 2014. Reverse polymerization of waste polystyrene through
747 microwave assisted pyrolysis. *Journal of Analytical and Applied Pyrolysis* 105, 35-42,
748 <https://doi.org/10.1016/j.jaap.2013.10.001>.

749 Victory, M., 2020. *INEOS, Plastic Energy to build pyrolysis-based chemical recycling plant*.

750 Williams, P.T., Williams, E.A., 2010. Product Composition from the Fast Pyrolysis of Polystyrene.
751 *Environmental Technology* 20, 1109-1118, <https://doi.org/10.1080/09593332008616908>.

752 Yang, M., Shibasaki, Y., 1998. Mechanisms of thermal degradation of polystyrene,
753 polymethacrylonitrile, and their copolymers on flash pyrolysis. *Journal of Polymer Science Part A:
754 Polymer Chemistry* 36, 2315-2330.

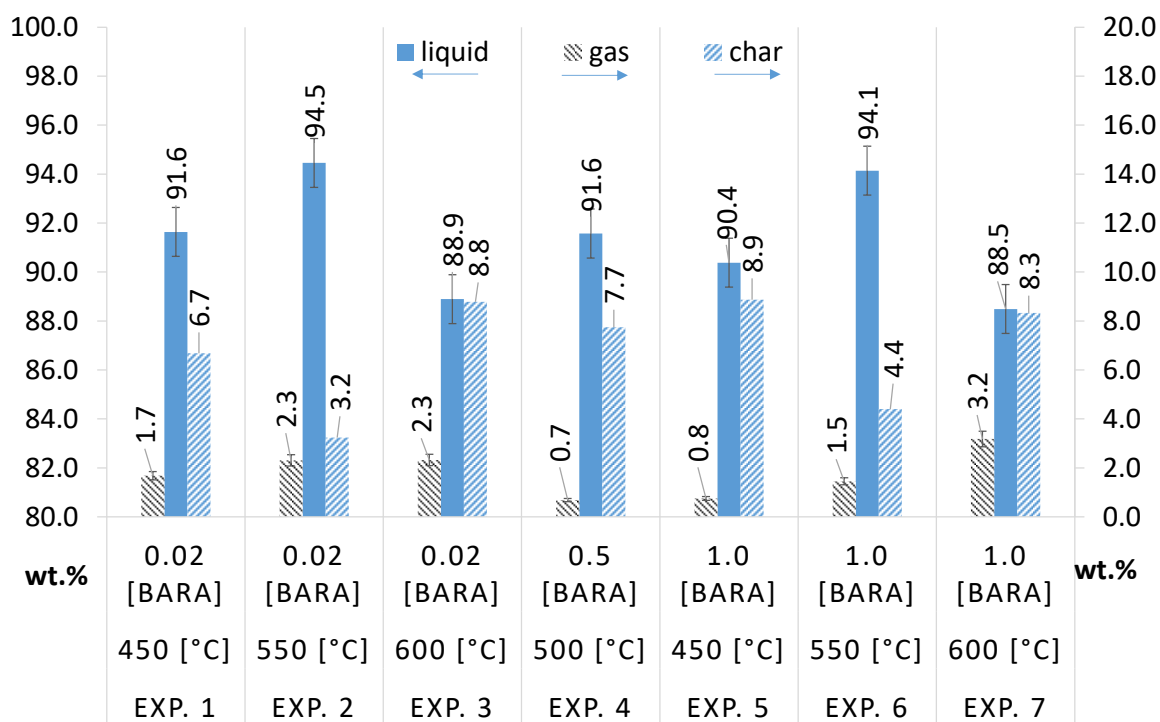
755 Figures and tables



756

757 *Figure 1 Flow diagram of the pyrolysis pilot unit: (1) extruder, (2) CSTR, (3) controlling cabinet, and*
 758 *PC, (4) N₂ cylinder, (5) N₂ volumetric flow controller, (6) condensers, (7) cooler, (8) back pressure*
 759 *controller, (9) vacuum pump, (10) off-gas flowmeter*

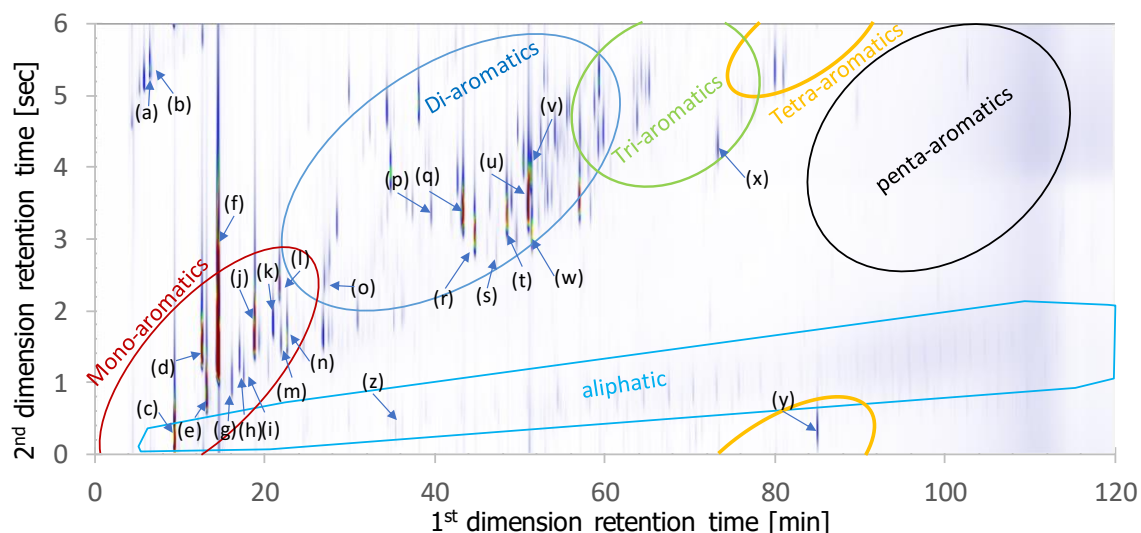
760



761

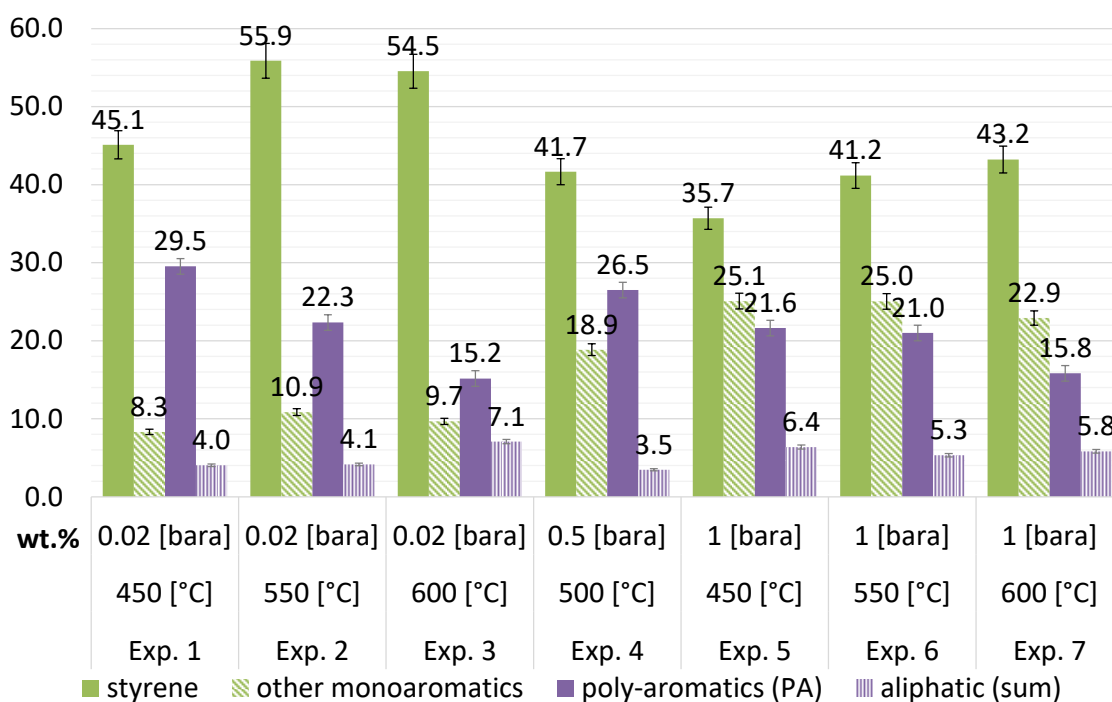
762

Figure 2 The mass balance of the continuous polystyrene pyrolysis in CSTR



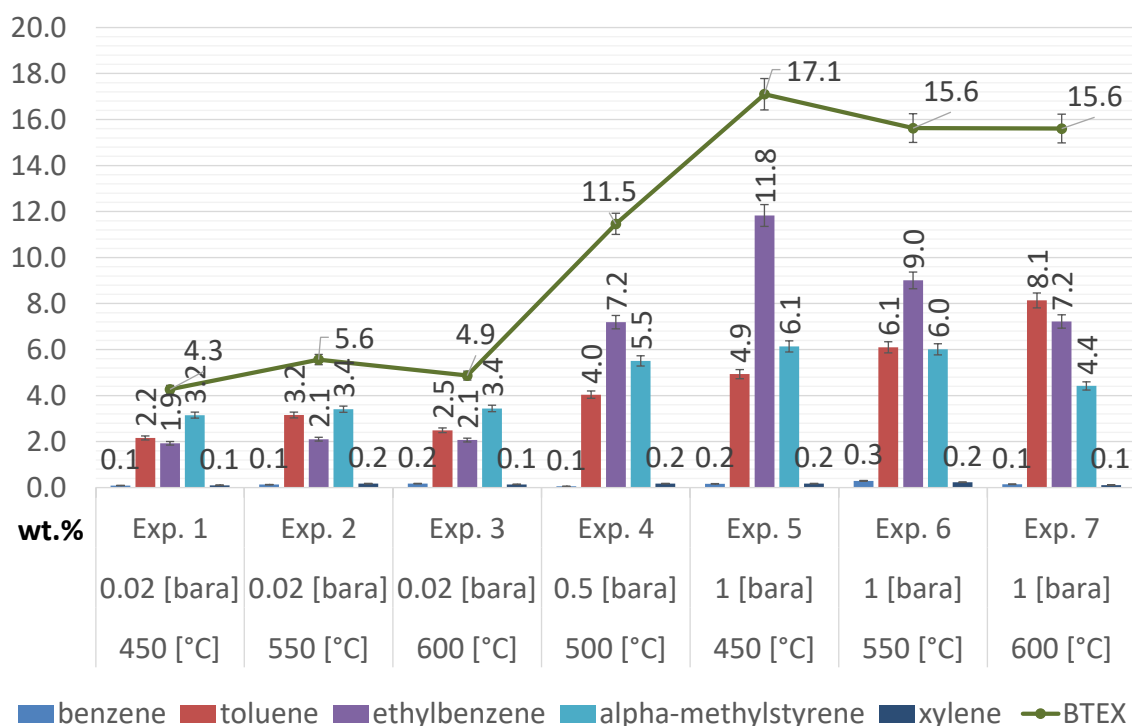
763

764 *Figure 3 GC x GC – FID chromatogram of the end-of-life polystyrene pyrolyzate with major*
 765 *compounds and group types. (a) Benzene (b) Cyclohexane (c) Toluene (d) IS=3-chlorothiophene (e)*
 766 *Ethylbenzene (f) Styrene (g) 1,2,4 trimethylbenzene (h) Benzene, 1-ethenyl-2-methyl- (i) Benzene,*
 767 *propyl- (j) Alpha-methylstyrene (k) Benzene, 1-ethenyl-2-methyl- (l) Indene (m) Benzene, 3-butenyl-*
 768 *(n) Benzene, (1-methylenepropyl) (o) Naphthalene (p) Diphenylmethane (q) Bibenzyl (r) 1,2-*
 769 *Diphenylpropane (s) 2,5-Diphenyl-2-hexene (t) 1,3-Diphenylpropane (u) 2,4-Diphenyl-1-butene (v) 1,3-*
 770 *Diphenylpropene (w) 2,4-Diphenyl-1-pentene (x) 2,4,6-triphenyl-1-hexene (y) 1,2,4-Triphenylbenzene*
 771 *(z) 2, 4, 6, 8-tetramethyl-1-undecene.*



772

773 *Figure 4 Effect of temperature (450, 500, 550 and 600 °C) and pressure (0.02, 0.5 and 1.0 bara) on the*
 774 *yield (wt.%) of the four main product groups found in polystyrene pyrolyzate*



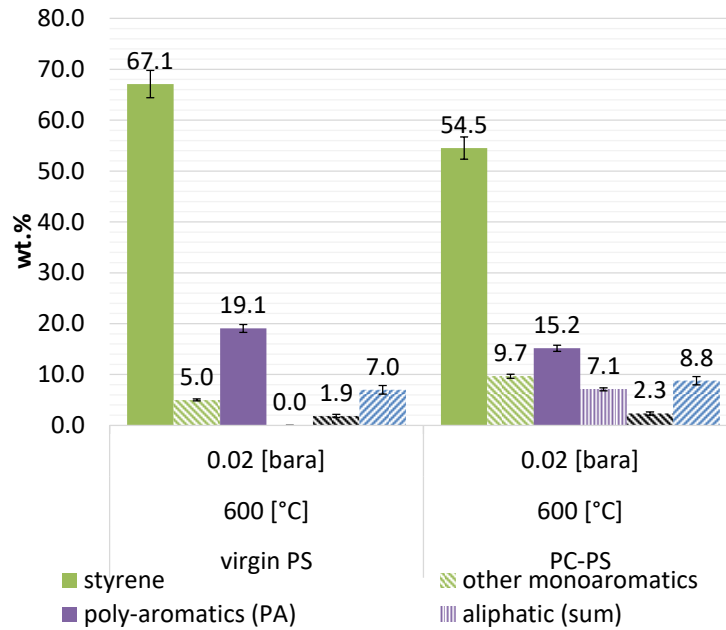
775

776 *Figure 5 Effect of temperature and pressure on the formation of BTEX and mono-aromatics excluding*
 777 *styrene.*

778 *Table 1 Yield (wt.%) of products from pyrolysis of end-of-life polystyrene (mono-aromatics (mon-A),*
 779 *di-aromatics (di-A), tri-aromatics tri-A), tetra-aromatics (tetra-A),)*

Experiment			1	2	3	4	5	6	7
P [bara]			0.02	0.02	0.02	0.5	1.00	1.00	1.00
T [°C]			450	550	600	500	450	550	600
Compound Name	Group type	C#	wt.% to the input feedstock						
gas (sum)	C1-C4		1.7	2.3	2.3	0.7	0.8	1.5	3.2
Liquid (sum)			91.6	94.5	88.9	91.6	90.4	94.1	88.5
toluene	mon-A	7	2.2	3.2	2.5	4.0	4.9	6.1	8.1
ethylbenzene	mon-A	8	1.9	2.1	2.1	7.2	11.8	9.0	7.2
styrene	mon-A	8	45.1	55.9	54.5	41.7	35.7	41.2	43.2
alpha-methylstyrene	mon-A	9	3.2	3.4	3.4	5.5	6.1	6.0	4.4
other mono.A	mon-A		1.1	2.2	1.7	2.1	2.2	3.9	3.1
1,3-Diphenylpropane	di-A	15	2.2	1.4	0.9	3.5	4.3	2.8	1.6
1,3-Diphenylpropene	di-A	15	0.4	0.5	0.5	0.9	0.9	0.7	0.5
2,4-Diphenyl-1-butene									
(dimer)	di-A	16	3.8	3.3	3.7	2.4	1.3	1.0	0.4
other di-A	di-A		4.5	9.2	8.5	7.9	6.4	10.0	9.8
C24H24 tri-A	tri-A	24	1.6	1.3	0.3	1.7	0.9	1.3	0.4
2,4,6-triphenyl-1-hexene									
(trimer)	tri-A	24	12.8	1.8	0.1	1.9	2.4	0.1	0.9
C25H26 tri-A	tri-A	25	0.7	1.2	0.1	1.4	0.9	0.7	0.2

other tri-A	tri-A	3.1	2.0	0.5	2.6	1.8	2.6	1.2
other tetra-A	tetra-A	0.5	1.7	0.5	4.0	2.6	1.7	0.9
aliphatic (sum)	aliphatic	4.0	4.1	7.1	3.5	6.4	5.3	5.8
others	others	4.7	1.2	2.5	1.1	1.6	1.6	0.7
char		6.7	3.2	8.8	7.7	8.9	4.4	8.3
total (yield to the input)		100.0	100.0	100.0	100.0	100.0	100.0	100.0



780

781

Figure 6 Comparison of product distribution using virgin PS and end-of-life PS as feed at pyrolysis conditions of 600 °C and 0.02 bara

782

783

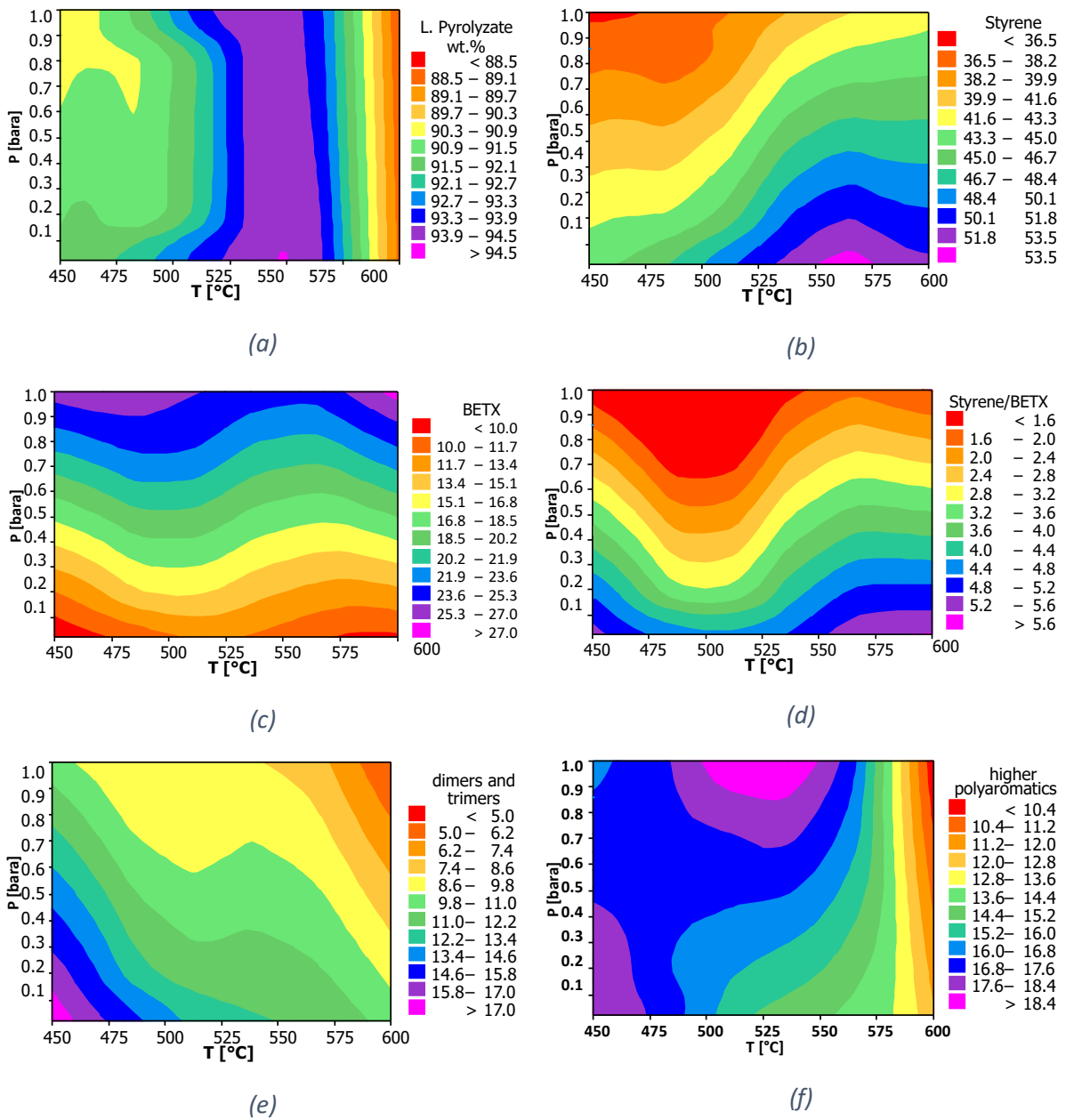


Figure 7 Product yield contour plot [wt.%] versus pressure [bara] and temperature [°C], **(a)** The contour plot of liquid pyrolyzate [wt.%] versus p and T , **(b)** The contour plot of styrene yield versus P and T , **(c)** The contour plot of (BTEX) yield [wt.%] versus P and T , **(d)** The contour plot of the ratio of styrene to BTEX versus P and T , **(e)** The contour plot of dimers and trimers [wt.%] versus P and T , **(f)** The contour plot of higher poly-aromatics [wt.%] versus P and T