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Best practice for sampling in automated parallel synthesizers†

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The ongoing digitalization leads to a higher degree of automation in chemical and material research. Particularly important is the sampling of reactions in order to monitor reaction kinetics and to enable online- and offline characterization. The sampling process must be well designed to take full advantage of the automated method without compromising analytical results compared to manual sampling. We present a new method for taking samples in an automated fashion for the application in synthesis robots and liquid handling robots obtained by simple additive manufacturing of a vial holder. This new method is systematically compared to standard approaches, which are currently the state-of-the-art techniques. The major benefit of the new technology is particularly important for easily evaporable solvents.

Introduction

The ongoing digitalization transforms chemical and material research in a significant manner.^{1,2} It enables fast screening for ideal reaction conditions or for the best performing material for a certain application.³ For this purpose, methods in the field of machine-learning and artificial intelligence are highly interesting.⁴ However, these methods require a large amount of data to be trained with and for this purpose, robot-based chemistry and combinatorial or high-throughput experimentation (HTE) are of particular interest.⁵ These methods enable the performance of a multitude of experiments simultaneously and fully automated.^{6,7} In addition, the high degree of reproducibility is an ideal precondition for the production of data for machine-learning algorithms.

Combinatorial and high-throughput experimentation were originally utilized in the field of pharmaceutical chemistry in order to replace or to complement the tedious search for new active compounds.⁸ Over the years, the new techniques found their way into further science fields such as polymer research⁹⁻¹¹ or catalyst screening.¹²⁻¹⁴ In general, HTE can help to explore a large data space in less time and without human interaction, reducing reproducibility issues.¹⁵ The main techniques utilized in this field are flow-based^{16,17} or robot-based systems using

However, in both systems, sampling is of great importance for the respective reactions in order to monitor reaction kinetics and to obtain a more detailed insight to reaction mechanisms. In general, there are several ways to analyze reactions. For flow applications, analysis can be either performed in-line, 20-22 where a probe is inserted into the flow and the measurement is performed in the process stream, on-line (a side stream is generated and analysis is performed in this stream), 23 at-line (a sample is taken from the flow stream and analyzed alongside the stream), 44 or off-line 22 (the sample is taken from the stream but processed elsewhere). 22,25 Several techniques are currently available to directly monitor the reaction progress in flow applications such as online IR spectroscopy, 22 nuclear magnetic resonance (NMR) spectroscopy or size exclusion chromatography (SEC). 27,28

For automated parallel synthesizers, however, online monitoring is more challenging as the reactions are performed in a batch-like manner. Hence, sampling in these machines is mostly performed off-line. Therefore, the robot's ability to collect liquid samples either in sampling vials or well plates is utilized. ^{19,29} Furthermore, there are few examples of implementations of direct injection of samples into an analytical device next to the synthesizer platform. ^{30,31} In case of sampling and direct insertion into the analytical device, it can be guaranteed that the sample leaving the reactor and the sample entering the analytical device are equal. In case of sampling into intermittent vials or other vessels, which is most prominent, a change of concentration or chemical composition cannot be ruled out.

For a perfect process in HTE, human intervention, such as vial capping, should be avoided to realize the full potential of this method. This approach would also utilize the possibility to

automated parallel synthesizers which are operating in a batch-like manner. 6,18,19

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† Electronic supplementary information (ESI) available: 3D models and GCode of

module: DSI. Videos of certain stages of the process: VSI (linked in the text). Setup of the evaporation experiments, overviews for evaporation experiments with different liquids, GC retention experiment and different views of the downholding module with dimensions. NMR and GC spectra figures. See DOI: https://doi.org/10.1039/d3dd00074e

take samples overnight, when no human is available to close the vials. Nevertheless, this is not possible with the aforementioned sampling method since sample components can evaporate from the vials over time changing the composition. While this is not a problem for (qualitative) methods analyzing the chemical structure of the product/educt, such as SEC, where the molar mass of the polymer is analyzed, it is a significant challenge for quantitative analysis, e.g., for kinetic determinations with mixtures of reactant, solvent and standards.29 Therefore, a thorough review of the sampling process is required, as it can limit the capabilities of automated parallel synthesizers for the kinetic investigations of easily evaporable substances. Consequently, we present in this article a completely new approach for quantitative sampling in automated research and compare this method with currently available state-of-the-art methods for parallel synthesizers to ensure the best retention for easily evaporable sample components.

Results and discussion

Concept of the study

The aim of the presented work is to consider the probable losses by evaporation for the sampling of easily evaporable liquids in automated parallel synthesizers and to identify a best practice for the sampling when such substances are present. To this end, first a short overview of the different configuration options for sample vessels is provided with respect to volume, material and capping options of the respective vessels. Subsequently, a selection of options is examined regarding their possible utilization for experiments with required sampling. For this purpose, different evaporation tests are conducted and practical considerations with regard to occurring spilling and required counter pressure utilization are made.

Configuration options for sample vessels

Generally, different influencing factors have to be considered when dealing with the sampling for easily evaporable liquids, thus, a short overview of the different configuration options for sample vessels is provided with respect to volume, material and capping options of the respective vessels.

Options for vials. A selection of possible sampling vessels is shown in Fig. 1. For vials, focussing on two main factors which influence the choice of the respective vessel for sampling seems to be appropriate. The first factor is the size of the vial. For this choice it is important to know the sampling challenge which should be solved. There are vessels in a range from only some hundred microliters to several dozens of milliliters or even laboratory flasks with liters of internal volume. For most of the screening applications, however, sample sizes (including solvents for quenching or subsequent analysis) are in the range of some hundred microliters to 2 mL.^{7,19} Hence, to avoid material as well as waste of space and to make a direct transfer into commonly used autosamplers possible, the best choice for a vial regarding volume is to use a 2 mL vial like shown in Fig. 1c.

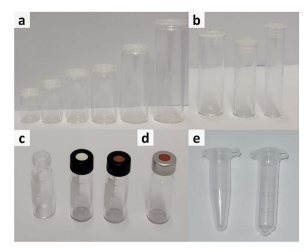


Fig. 1 Overview of several utilizable vial options. (a) Flat bottom snap cap vials in different internal volumes from 3 to 50 mL. (b) Flat bottom vial with polyethylene plug cap (1 to 2 mL). (c) Screw neck autosampler vials with screw caps (2 mL). (d) Crimp neck autosampler vial with crimp septum cap (2 mL). (e) Polypropylene tubes wit plug cap (2 mL).

Another important factor is the choice of material. The most common materials chosen for production of vessels are either polypropylene (Fig. 1e) or glass (Fig. 1a-d). Depending on the conducted experiments this choice might change due to chemical incompatibilities or other necessities. However, for most experiments, glass vials are the best option as the material is chemical inert, stable to a range of temperatures and vials are available in a great range of volumes.

Options for lids. A further factor potentially affecting the efficacy of sampling is the sealing of the vial. The possibilities range from snap-on lids and plugs (Fig. 1a, b, and e) over screw caps (Fig. 1c) to crimp caps (Fig. 1d). If used in automated parallel synthesizers and autosamplers, it is important to gain the possibility to sample through the lid of the vial. This is only the case for lids containing septa, as the other materials such as the polyethylene of the snap-on lids are too rigid to be pierced by the needle of the automated parallel synthesizer. Another factor is the possibility to remove the cap after the sample preparation for further processing of the sample, if required. For this purpose, screw on as well as snap cap or plug vials are better suited in comparison to crimp vials, as the removal of the lid and, thus, the accessibility of the sample is more facile.

Considering all previously stated arguments, the best choice for sampling are 2 mL glass vials with, if required, a screw top lid containing a septum. Hence, this combination was chosen for further investigation.

Test for the requirement of sealing

After a first determination of the most suited vial option, further investigations were performed regarding the necessity for closing the vial during sampling and the best lid option. The available different sealing options are shown in Fig. 2 and comprise of either an unsealed vial or a vial sealed with a septum cap.



Fig. 2 Overview of sealing options for the subsequent experiments. Left: unsealed vial. Middle: vial with slit septum lid. Right: vial with regular septum lid.

The easiest way to sample would be to use an unsealed vial as the omission of a cap is economically as well as ecologically advantageous. However, it is probable that for easily evaporable substances evaporation occurs which could change the sample composition. To test the hypothesis of sample evaporation from open vials in the first place, a simple evaporation test was performed with a solution of a dye in a low boiling solvent (bromophenol blue in methanol (boiling point: 65 °C (ref. 32))). Two experimental rows (E1 and E2) were conducted. For E1 the vials were capped with a septum cap, while for E2 no sealing was performed. The results of the experiments are shown in Fig. 3. The degree of filling for the two experiments varies significantly over time. For E1 no evaporation occurs. The filling level of the

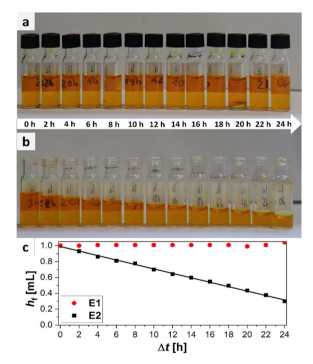


Fig. 3 Overview of results of the evaporation experiment. (a) Picture of sealed vials for different elapsed times since filling (E1). (b) Picture of unsealed vials filled at the same time as the sealed ones (E2). (c) Plot of filling height $h_{\rm f}$ of sealed (red) and unsealed vials (black), against the elapsed time between evaluation and filling Δt .

vials stays nearly constant in the range of accuracy of the robotic liquid dispensing system and is not dependent on the time of filling. This can be seen, as the filling height of the sample taken in the beginning (24 h elapsed between picture and filling) (E1/24 h) is the same as for the sample taken after 24 h (E1/0 h). For E2, however, a steady decrease can be observed from 100% filling height (1 mL) for E2/0 h to the primarily filled sample E2/24 h where only about 0.29 mL remain inside the vial. This linear decrease is also visible in the plot (Fig. 3c). With the data from the plot, an empirical evaporation rate for methanol in this context can be determined according to the slope of the linear fit in Fig. 3c, which is:

$$h_{\rm f} = 0.989 \text{ mL} - 0.028 \text{ mL h}^{-1} \cdot \Delta t$$

Hence, the evaporation rate of methanol for this experiment is 28 μ L h $^{-1}$. According to this information, 1 mL of methanol would evaporate in ca. 36 h in an open standard autosampler vial under the given conditions. If methanol (1 mL) is used as standard substance and a sample is automatically drawn by the parallel synthesizer at 8 p.m., the result of the quantitative measurement, performed at the next day, when the researcher appears in the laboratory at 8 a.m., will already be flawed by approximately 34%. In summary it is obvious that evaporation occurs in parallel synthesizers when sampling vials are left open for low to medium boiling substances. As evaporation did not occur for the samples stored in capped vials as obvious from samples E1/0 h to E1/24 h, those should be used for sampling instead.

Dependence of evaporation on vapor pressure of liquid

An influential factor for the evaporation of liquids is the vapor pressure of the substance. To claim the best sampling practice, it is important to monitor the sampling over a larger area of different liquids with differing vapor pressures. Hence, the evaporation behavior of six common liquids with a vapor pressure difference at 20 °C of *ca.* 580 mbar was investigated. The chemicals with their respective vapor pressures and experimentally determined evaporation rates are shown in Table 1.

For the experiment a similar setup was utilized as for the experiment of the requirement of sealing. The only difference being the storage of the liquids before the filling of the sampling vials.

The figures with an overview of the vials and the results of the evaporation experiments and plots of the evaporation rates are presented in the ESI (Fig. S3–S8†).

As visible from Table 1, overall evaporation in the investigated system can firstly be monitored above vapor pressures of 130 mbar. For lower evaporating liquids like dimethyl formamide (L1) or water (L2), no evaporation can be observed in neither case (open or sealed vial) during the time of observation (24 h). For more facile evaporating substances like methanol or acetone with vapor pressures above 130 or 200 mbar, evaporation can be observed from the unsealed vials L30 and L40 with rates between 34 to 81 μ L h⁻¹. This would correspond to total evaporation times of 1 mL of sample between *ca.* 30 (L3) and 12

Table 1 Overview of the dependence of the evaporation on the vapor pressure with literature known vapor pressures and empirically determined evaporation rates for different liquids

Experiment number	Liquid	Vapor pressure @ 20 °C [mbar]	Evaporation rate [mL h ⁻¹]	
			Open vial o	Sealed vial s
L1	Dimethylformamide	5.33 33	0	0
L2	Water	23.38 34	0	0
L3	Methanol	130.11 35	-0.034	0
L4	Acetone	246.38 ³⁶	-0.081	0
L5	Dichloromethane	481.29^{37}	-0.166	-0.005
L6	Diethyl ether	583.55 ³⁸	-0.500	-0.015

hours (L4). In this range of vapor pressures, however, the sealed vials L3s and L4s show no evaporation and, hence, keep their initial composition over a time of at least 24 h. With further increasing vapor pressure, the challenge of absolute retention becomes greater. As the system using a slit septum lid is only semi-closed, increasing build-up pressures inside the vial due to evaporation of the liquid in the vial's headspace are able to be released. This can be observed for dichloromethane (L5) and diethyl ether (L6). For both, in case of the unsealed vials, total evaporation occurred within 24 h. For L6o total evaporation already occurred after 2 h, while for L50 the same was the case after ca. 6 h. For the sealed vials, as well, evaporation could be observed. However, the rates of evaporation were decreased significantly compared to the unsealed vials. The decreasing factor, determined by dividing the evaporation rate for the open vial by the rate obtained for the closed vial is ca. 33, which would mean that a sample which evaporates within one hour in an opened vial would need notably more time (33 h) to evaporate completely from a sealed vial under similar circumstances.

Generally, it is obvious that with increasing vapor pressures, the task of sealing is getting more difficult and deviations between the initial composition at the time of sampling and the time of measurement become more probable. For liquids below the vapor pressure of water, no visible advantage could be found for the proposed method as no evaporation occurs even from the unsealed vials (during the observed time frame of 24 h). However, a lot of used chemicals, e.g., monomers in polymerizations or solvents, have vapor pressures above the ones of the mentioned substances.39,40 Hence, the area of higher vapor pressures is of greater interest. With increasing vapor pressure, the proposed method shows its advantages in comparison to the classic use of unsealed vials. The rate of evaporation decreases significantly when using sealed vials which, in turn, leads to a better retention of the sample's properties between the time of sampling and the time of measuring.

Dependence of evaporation on external factors

Besides the previously investigated parameters, the evaporation from open and closed vials is dependent on further factors. The first group are environmental factors, like humidity or surrounding temperature. These are dependent on the setup used and the experimental approach. In the used experimental setup, the heat radiated from the reactors in close proximity to

the sampling racks plays a vital role. *E.g.*, for water the evaporation will be faster if the temperature is increased or the surrounding relative humidity is decreased.⁴¹ In this experiment, the values for the temperature, measured close to the sampling rack, reached values of up to 29.3 °C at the end of the interval of 24 h. The relative humidity, in turn, decreased by *ca.* 8% from 44.3% to 36.5% over the course of the experiment. Differences in these temperatures and humidities will definitely affect the evaporation of samples taken in non-sealed vials. For sealed vials, the influence will be smaller as the system is semiclosed. However, in general, the changes in temperature or humidity inside the automated platform will not be large but rather small, even with higher heating temperatures of the reactors. Hence, the assumptions taken from the evaluation of the conducted experiments are generalizable.

Additional influential factors could be the specifications of the needle. In the case of the performed experiments, a septum piercing needle with an outer diameter of 1.5 mm was utilized. This will, in turn, lead to a small puncture through the septum material and low exchange between the atmosphere in the vial and outside it as the physical barrier remains intact. With larger needle diameters the lid material could be pierced in a way that the material is irrecoverably damaged. The resulting holes would then lead to easier evaporation. Therefore, it is recommended to take the needle with the smallest applicable diameter.

Furthermore, the surface area will play an important role for evaporation. The higher the area, the larger also the evaporation tendency.⁴² However, as previously mentioned, the best choice are small (2 mL) vials which in turn always have the same surface area. Hence, this point will not be investigated in detail in this study.

Comparison of sealing options

As mentioned earlier, there are two different available septum types (see Fig. 2). The first consists of a silicone/PTFE septum with a slit in its center. The second option is the usage of an unslitted rubber/PTFE septum. In the following, a detailed comparison for their application is provided.

Reaction mixture evaporation. To examine if the caps differ in terms of their sealing abilities, the effect of the evaporation on a mixture for a typical polymerization was investigated. An attempt was made to evaluate the effect of evaporation on the

preservation of the ratio between a monomer (methyl methacrylate (MMA)) and a standard substance (anisole) in tetrahydrofurane (THF). For this purpose, seven individual samples were prepared. Each of the samples contained 75 µL of the described mixture. A first sample (V1) was prepared by adding 0.7 mL of deuterated chloroform and sealing the vial with a regular rubber septum cap. This reference sample was then stored at 5 °C for the course of the experiment. The remaining six vials, containing 75 µL of sample, were separated into two groups of three samples. For the first group (V2) the 75 µL of sample remained in the vial and no further addition of substances was performed. For the second group (V3), 0.7 mL of deuterated chloroform were added to the sample in the vial. Afterwards, one of the vials of each group was closed with a regular rubber septum lid (V2/3c). Another was closed with a slit septum cap (V2/3s), while the third one remained unsealed (V2/30). Subsequently, the samples were stored for 24 h in a sampling rack next to a reactor block, which was heated to 70 ° C. For each of the samples a ¹H-NMR spectrum was recorded (see Fig. S9-S14†). The integral of the *ortho* and *para* protons of anisole ($\delta = 6.75$ to 6.95 ppm) was calibrated to 3 and the according methylene proton integral for MMA ($\delta = 6.02$ ppm) was revealed (see Fig. S9†). This integral was then compared to the one from V1 to understand in which dimension the evaporation had altered the sample composition. The results of these calculations are represented in Table 2. Furthermore, an overview of the respective NMR spectra for samples with preadded deuterated chloroform is provided in Fig. 4. As it can be seen from the table and figure, the samples in the unsealed vials evaporated completely, which is also consistent with the prior test. For the four sealed combinations, higher ratio retention was achieved for the combinations with pre-added chloroform (V3c and V3s). For those samples, a complete retention was achieved. For the two non-topped-up samples (V2c and V2s), evaporation of the sample occurred and the ratio changed due to a higher evaporation rate of MMA compared to anisole.

This effect is more pronounced for the combination with the slitted septum. However, both combinations without prefilled solvent cannot be used for an error-free sampling strategy. Hence, it is required to store the samples prior to analysis mixed with a larger volume of solvent or quenching agent as otherwise

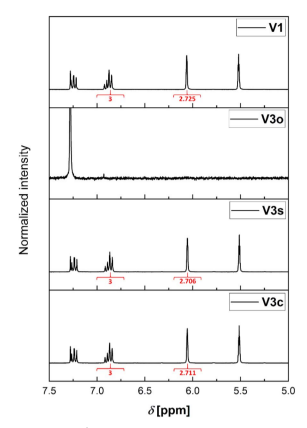


Fig. 4 Zoomed in ¹H-NMR spectra of the samples with pre-added deuterated chloroform with indicated integrals, where applicable From top to bottom: reference sample (stored at 5 °C). Unsealed (open) vial. Slit septum lid. Regular septum lid.

the changes in the ratio of the components of the sample are severe.

This data is also in accordance with the conducted sampling experiment for GC measurements. The data for this experiment (V4, V5) can be found in the electronical ESI (Table S1†) of this

Dependence of evaporation on number of septum punctures. Since individual vials could also be utilized as reaction vessels in the future, it is of interest to examine whether both sealing options behave equal with respect to multiple punctures

Table 2 Retention of ratio of integrals for different sampling conditions in comparison to a reference sample stored at 5 °C over the course of the experiment

Sample condition	Experiment	Integral for MMA methylene protons a	Retention of ratio [%]
Reference sample	V1	2.725	100
Open vial, no p-chloroform addition	V2o	<i>b</i>	<i>b</i>
Open vial, D-chloroform addition	V3o	<i>b</i>	<i>b</i>
Regular septum lid, no D-chloroform addition	V2c	0.735	27
Regular septum lid, p-chloroform addition	V3c	2.711	100
Slit septum lid, p-chloroform addition	V2s	0.181	7
Regular septum lid, no D-chloroform addition	V3s	2.706	100

^a Integral of the peak at $\delta = 5.52$ ppm determined compared to the integral (3) of the *ortho* and *para* anisole protons ($\delta = 6.75$ to 6.95 ppm).

^b Complete evaporation of the sample. Hence, no integral and no retention measurable.

through the septum. Therefore, two evaporation experiments were performed to investigate the effect of multiple punctures on the different lid types and the corresponding evaporation. The utilized solvent was either methanol (P1) or dichloromethane (P2), since both are standard solvents for chemical reactions and feature different vapor pressures (see above). The two different sealing options, a slit septum lid (abbreviation sl) or a regular rubber septum lid (abbreviation rl) were utilized. The experimental setup was similar to the already applied one for the investigation to determine the dependence of the evaporation on the vapor pressure. However, the vials were prefilled with 1 mL of solvent solution and then punctured multiple times (from 0 (e.g., P1/sl/0) to 30 times (e.g., P2/rl/30)). The maximum number of 30 punctures was chosen with respect to the internal volume (2 mL) of the selected sampling vials. The overview of the observations for all vials and their filling level are presented in the ESI (Fig. S22 and S23†).

As it can be seen from the mentioned figures, no direct correlation can be obtained between the number of punctures and the evaporation tendency of the liquid inside the vial. The evaporation seems to be almost identical for most of the vials for both of the sealing possibilities. This is particularly evident for the experiment with methanol P1 (Fig. S22†). In this case, all vials are filled to the same level after the experiment, regardless of the number of punctures or the lid type itself. For P2 (Fig. S23†), however, smaller differences are visible. Nonetheless, there does not seem to be a clear tendency which sealing type works better. For the samples with zero punctures, the regular sealing lid retains more liquid. This is consistent with expectations, as the slit in the septum dissipates the excess pressure which builds up for liquids with high vapor pressure (dichloromethane in this case) under the experimental conditions. The regular septum lid does not provide this capability, which is why it performs better (see Fig. 5). For the samples with ten, 14, 22 and 28 punctures, the slit septum cap seems to be the better choice as it retains more liquid. For the other puncture numbers, there is not much difference. As the difference is very pronounced for the samples with 28 punctures, these were investigated further and are shown in comparison with the samples with zero and 30 punctures in Fig. 5. For the vials with 28 punctures (P2/rl/28; P2/sl/28) through the lid, it can be seen that the lid of the vial sealed with a regular rubber septum lid (P2/rl/28), was damaged to such an extent that a crack formed, resulting in total evaporation of the dichloromethane. Since this phenomenon did not occur for P2/rl/30, it can be assumed that the number of punctures is less important than the location of the recurring punctures. This depends on the accuracy of the placement of the needles within the automated platform and varies depending on the calibration state of the X- and Yaxis of the platform and the platform itself. The slit septum lid has an advantage in this regard, as the pre-existing slit provides a "guide" for the needle. The regular rubber septum does not offer this possibility. Therefore, it is more likely to be punctured at different locations. This can cause a tear to form in the septum since the following punctures are close to the previous puncture. Should such a tear occur, it will result in faster evaporation of liquids from the vial sealed with a regular rubber

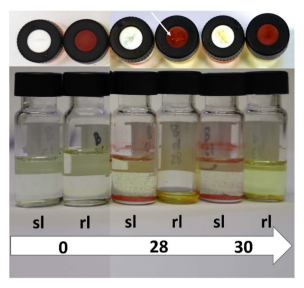


Fig. 5 Overview of the lids with zero, 28 and 30 punctures of experiment P2 with the respective filling level of the vials after the experiment. The arrow points to the tear in the septum.

septum lid. However, the occurrence of such a crack seems to be at least partially random, as shown by the comparison of P2/rl/28 and P2/rl/30.

Choice of sealing option. As apparent from the performed experiments, sealing of the vials is required in the context of reaction sampling. There are different sealing options as presented in Fig. 2. Since both options (regular rubber/PTFE septum cap and slit silicone/PTFE septum lid) are utilizable in the context of ratio retention (with prefilled quenching agent), other parameters determine the choice of the ideal sealing solution. Initial reaction tests were performed using the regular septum lids as they were readily available in our group's laboratories. However, during the course of sampling it became apparent that spillage was occurring. As a consequence, the sampling process was examined in more detail. The immanent steps of the subsequent test, utilizing a prefilled vial with regular rubber septum cap and a vial with slit septum cap are presented in Fig. 6. Apparent from Fig. 6b, the spillage occurred at the moment the needle was withdrawn through the regular rubber septum cap, while this phenomenon could not be observed for the slit septum cap at the same stage of the process (Fig. 6f). A corresponding video of the process for the regular rubber septum cap can be followed in the ESI (VSI1†). Due to the prefilling of quenching agent and the subsequent waiting time, a pressure seems to have built up inside the vial. This excess pressure cannot escape in the case of the regular rubber septum lid. During the injection of the sample, the pressure is further increasing. As the needle is retracted from the vial (Fig. 6a), a small hole is created in the septum and the overpressure is released (Fig. 6b and c) by leaking out of the vial. In addition, the pressure causes a problem with the injection needle. The needle is connected to a syringe pump (piston pump). The pump draws up the sample by creating a negative pressure inside the syringe through movement of the piston. When the

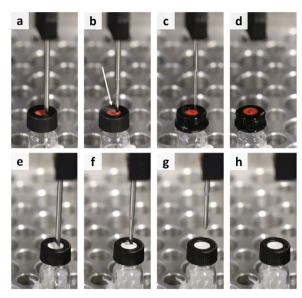


Fig. 6 Overview of the needle retraction process for the vials with regular rubber septum (a-d) and slitted rubber septum (e-h) at the same steps in the process.

sample is dispensed into a vessel, the plunger moves upward and creates an overpressure that drives the liquid out of the connected tubing and needle. However, this expulsion is hindered by an overpressure in the target vessel. Hence, not all of the sample volume is injected into the vial leading to errors for the sampling. As soon as the needle is withdrawn from the vial featuring overpressure, the pressure generated by the syringe pump is released and a part of the sample is spilled outside the vial (Fig. 6c). This poses a problem for reproducible sampling, since sample gets lost during the process, which is why the regular rubber septum cannot be utilized in a suitable manner. One alternative would be to use vials without lids for sample transfer and to cap them immediately afterwards to avoid pressure build-up and accordingly spillage on the one hand and evaporation on the other. However, this would reduce the advantage of the currently available automated systems, as human interaction would be required.

Another option is to use caps with a slitted septum as they possess the ability to release the excess pressure before the sampling can get affected (Fig. 6e-h). Furthermore, excessive evaporation is also avoided. Moreover, in this manner no human intervention is required, which maintains the advantage of automated platforms to work without manual intervention.

Practical consideration for sampling - downholding

At this point, the best sampling practice seemed to be determined by utilizing a 2 mL glass autosampler vial with a screw-on slitted septum lid. However, when applying the sampling method in the automated parallel synthesizer, another challenge arose. As shown in Fig. 7, withdrawing the needle from the vial caused the vial to be pulled out of the sample rack. As tested, this occurs regardless of the caps used. This pulling up of the vial results in loss of the vial within the robotic platform, as stripping the vial from the 4-needle head (Fig. 7c) causes the

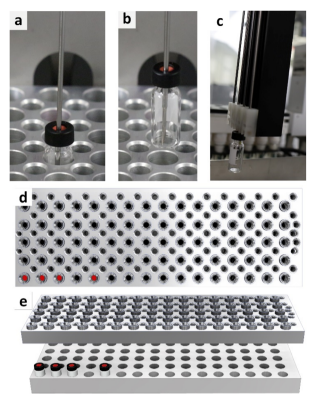


Fig. 7 Representation of a vial, withdrawn with the needle of the 4needle head (a-c) and representation of the developed downholding module in top view (d) and orthogonal view (e).

vial to fall at undefined places in the robot. A video of this process can be accessed via the ESI (VSI2†). This challenge could be overcome by utilizing fused deposition modelling 3Dprinting. Therefore, a module to hold down the vial was designed (design files available in the ESI (DSI1-DSI3†)), printed, and tested in the context of applying counter-pressure onto the vial. The module applied on top of a sample rack with four vials is represented in Fig. 7d and e. A video of the module put into work (VSI3) as well as a video of the module with integrated vials on top of a sampling rack are part of the ESI (VSI4†). The module was able to keep the vials in place during the sampling process and was, therefore, applied for subsequent experiments.

Final sampling practice

The finally applied and recommended sampling strategy is the utilization of a prefilled 2 mL glass autosampler sampling vial with a slit septum cap to avoid ratio change. Furthermore, a module for downholding of the vial during the sampling process should be applied. With this method, a reproducible sampling can be achieved. Furthermore, this method can be utilized also for highly volatile substances to reach less evaporation from the vial. The method is easily scalable as only the number of vials needs to be increased and more downholding modules need to be manufactured. This is fairly easy as filament 3D-printing nowadays is a standard method in laboratories and hobbyists workshops.

Experimental

Materials and methods

Methyl methacrylate was purchased from Sigma-Aldrich in 99% purity. Bromophenol blue was purchased from Merck. Chloroform was purchased in HPLC-grade from VWR. Deuterated chloroform was purchased from Eurisotop. Dimethylformamide (>99.9%), acetone (>99%), dichloromethane (98%) and diethyl ether (technical grade) were purchased from VWR chemicals. Methanol (>95%) was purchased from Thermo Fisher Scientific. Water was taken as tab water from the laboratories fresh water system.

Humidity and temperature in the sampling rack were measured with a Lufft OPUS20 TCO.

 1 H NMR spectra were recorded on a Bruker Avance I spectrometer (300 MHz) at 298 K. The chemical shifts are given in parts per million (ppm on δ scale) related to the deuterated solvent.

GC measurements were performed on the following system: Shimadzu GC-2010 with an FID (detector), Carl Roth, Roti®Cap-5 MS (30 m long, 0.25 mm ID, 0.25 μ m film thickness) (column) with a stationary phase consisting of (5% diphenyl) 95% dimethyl polysiloxane and helium as carrier gas. The samples were prepared in chloroform as solvent.

All experiments/tests were conducted utilizing a Chemspeed Accelerator SLT 106 automated parallel synthesizer platform. The synthesizer was equipped with a 4-needle head for liquid transfers with connected 10 mL tubing and syringes, one reactor block consisting of 16 individual reactor vessels (13 mL) with heating mantles and a sample rack (double level holder, purchased from Chemspeed). The reactor block was topped up with a reflux module consisting of individual hollow glass fingers and a controlling module to open and close the reactors. Heating to 70 °C was supplied by an external heating circuit with a Huber Unistat Tango as a dynamic temperature control system. The reflux coolers were flowed through with a mixture of ethylene glycol and water (volumes = 1:5) to apply a constant cooling of the fingers to 5 °C. In this case a Lauda Microcool MC

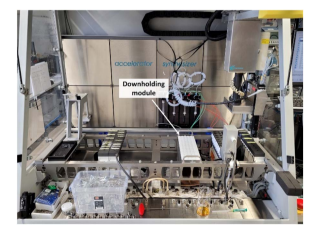


Fig. 8 Overview of the robotic platform with 4-needle head (right), syringe pumps (middle) as well as sample rack with white downholding module and reactor block (middle right).

600 was utilized as temperature control system. The platform is shown in Fig. 8.

The downholding module was designed in Autodesk Inventor Professional 2020 and printed on a Prusa i3 MK3S fused deposition modelling 3D-printer with polylactic acid (PLA) Filament (PrimaValue PLA, different colours, diameter = 1.75 mm) (GCode available as DSI3†).

Test for the requirement of sealing

The previously described synthesis platform was utilized. Four of the reactors were filled with 10 mL of bromophenol blue (BPB) in methanol ($c=2.99~{\rm mmol~L^{-1}}$). The solutions were heated to 70 °C and mixed via vortex-shaking at 400 rpm. The vials with a septum cap (E1) and without a lid (E2) were placed in a sample rack in a distance of about 11 cm from the reactors (see Fig. S2†) for 24 h. Every second hour 1 mL of sample was taken with the automated liquid handling system from two of the reactors and filled into the respective vials at the same time. After 24 h, the sampling rack was removed from the synthesizer and the filling level of each vial was documented.

Dependence on vapor pressure of liquid

The previously described synthesis platform was utilized. Two 100 mL vials were filled each with 75 mL of a solution of a solvent with 20 to 25 mg of BPB. For each solvent sampling vials with septum caps (*e.g.*, L1s) and without a lid (*e.g.*, L1o) were placed in a sampling rack about 11 cm from the reactors. Then, the reactors were heated to 70 °C at a vortex-shaking rate of 400 rpm. Every second hour 1 mL of sample per sampling vial was taken with the automated liquid handling system from the 100 mL stock vials and dispensed into the sampling vials. After 24 h the sampling rack was removed from the synthesizer and the filling level of each vial was documented.

Comparison of sealing options

Reaction mixture evaporation. A solution of 8.54 mL tetrahydrofurane, 0.4 mL (3.68 mmol) anisole and 1.06 mL (9.95 mmol) methyl methacrylate was prepared. 75 μL of sample were taken from the mixture and transferred into eight 2 mL screwtop vials. For four vials, the samples were topped up with 0.7 mL of HPLC-grade chloroform. A topped-up sample (V1) was stored in the refrigerator at 5 °C until the end of the experiment. Additionally, the remaining six vials were paired so that a vial with topped up solvent (V3) could always be compared to a nontopped-up one (V2). One of these pairs remained unsealed (V2o and V3o). Another pair was closed with a regular screw-on rubber septum cap (V2c and V3c), while the last pair was sealed with a slitted ruber septum cap (V2s and V3s). For better comparability with actual experiments, the septa were pierced three times with a needle to simulate the sampling process inside an automated parallel synthesizer. Afterwards, the vials were placed in a sampling rack inside the automated parallel synthesizer according to the description for the first test. The reactor block next to the sampling rack was heated to 70 °C to simulate a polymerization reaction. After 20 h and 33 min, the samples were taken from the sampling rack and samples

without topped-up solvents were topped up with 0.7 mL of deuterated chloroform. Subsequently, ¹H-NMR measurements of all samples were performed.

Dependence of evaporation on number of septum punctures. The previously described synthesis platform was utilized. A solution of 70 mL of a solvent (either methanol (P1) or dichloromethane (P2)) with 20 to 25 mg of bromophenol blue was prepared. 32 Vials were filled with 1 mL solution each. 16 of these vials were sealed with a regular septum lid (e.g., P1/rl). The remaining 16 vials were sealed utilizing a slitted septum lid (e.g., P1/sl). The vials were placed inside a sampling rack. Afterwards, the lids were punctured utilizing a septum piercing needle from 0 (P1/sl/0; P2/rl/0) to 30 (P1/sl/30; P2/rl/30) times in steps of two punctures to model a repeated aspiration of reaction components from the vial. The sampling rack was placed inside the synthesis platform and the reactors next to the rack were heated to 70 °C at a vortex-shaking rate of 400 rpm. After 24 h the sampling rack was removed from the synthesizer and the filling level as well as the condition of the lid of each vial was documented.

Choice of sealing option. Three standard 2 mL glass auto-sampler screw cap vials (surface area: ca. 0.66 cm²) were placed inside a sampling rack. One of them was left unsealed, one was capped with a regular rubber septum cap and the last with a slit septum cap. To avoid retraction of the vials due to friction between the needle and the septum, the vials were wedged with paper into the respective well of the sample rack. Afterwards, 0.1 mL of chloroform was inserted to the vials using a septum-piercing needle with an outer diameter of 1.5 mm and a waiting time of approximately ten minutes was applied. Subsequently, 1 mL of chloroform was injected to the vials via the 4-needle head of the automated parallel synthesizer (Chemspeed SLT 100). After the injection, the needle was retracted from the vial.

Conclusions

High-throughput experimentation is important for the ongoing digitization of material sciences and chemistry in order to obtain reliable data for artificial intelligence and machine learning approaches. A special emphasis in this field should lie on the sampling process as this presents a crucial step and can affect the reaction data in a significant manner. In particular, quantitative sampling is a challenge in automated parallel synthesizers as evaporation from open vials can change the composition of a sample. To enhance this part of the automated process, we developed a new method for quantitative sampling. Hence, several options regarding the choice of the ideal sampling vessel as well as sealing option were discussed and investigated with regard to their suitability for sampling of easily evaporating substances. It was found that sealing of the sampling vials, in contrast to the current state-of-the-art, is of utmost importance to avoid complete or partial evaporation of the sample. For partial evaporation over time, it could be shown that the ratio of a particular monomer (MMA) changes with respect to a standard (anisole). Hence, sampling should be performed with prefilled quenching agents or solvents. The ideal sealing option for sampling could be found in a 2 mL

autosampler glass vial with a slitted septum due to its ability to release any excess pressure that occurs during injection of the sample into the vial. Finally, the use of a counter-pressure module was found to prevent the vial from being withdrawn from the sampling rack.

Therefore, it was possible to establish a new best practice for automated sampling for various analytical methods such as GC or SEC in automated parallel synthesizers. These findings will significantly improve the possibilities for representative quantitative sampling in synthesis robots and, therefore, open the possibility to generate more reliable data for the use with artificial intelligence methods which will foster the understanding of quantitative structure–property relationships.

The design files for the downholding module as well as the G-Code for 3D printing on a Prusa i3MK3S with polylactic acid filament can be found in the ESI (DSI1-DSI3†). ¹H-NMR as well as GC primary data can be obtained on request from the authors, the processed spectra are shown in the ESI.†

Data availability

The data, pictures, videos and design files supporting this article have been uploaded as part of the ESI.† Further raw data can be obtained by the authors on request.

Author contributions

Conceptualization: M. R. and S. Z.; investigation and data curation: M. R.; design of module: M. R. and T. S.; writing – original draft: M. R.; writing – review and editing: M. R., T. S., S. Z. and U. S. S.; supervision: S. Z. and U. S. S.; resources: U. S. S.; project administration: U. S. S.; funding acquisition: U. S. S. All authors have read and agreed to the published version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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