

# One pot/two donors/one diol give one differentiated trisaccharide: powerful evidence for reciprocal donor–acceptor selectivity (RDAS)

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Three component, one-pot reactions involving equimolar amounts of the acceptor diol and both armed and disarmed donors presented simultaneously, produce a *single* double-differential glycosidation product; this phenomenon provides evidence for Reciprocal Donor Acceptor Selectivity (RDAS).

The traditional protocol for differential, double-glycosidation of an acceptor diol requires a series of programmed protection/deprotection steps to ensure that only one of the acceptor-OHs is presented to each donor at each coupling event.<sup>1,2</sup> Thus, for the case illustrated in equation (i), a *minimum* of four steps would be needed to obtain compound **3** from diol **1**, without contamination of the diastereomer **4** (and avoidance of the symmetrical competitors **5** and **6**). The direct process, **1**→**3**, is generally thought to require exquisite regioselective finesse and therefore is best left to enzymatic procedures.<sup>3</sup>

However, in this manuscript, we report the development of differential, double-glycosidations of diols in which donors are virtually 'told' where to go, thereby enabling direct conversions of the type **1**→**3**, without contamination of regioisomer **4**.

We recently reported that a variety of diols, including **7**, **8** and **9**, underwent regioselective glycosidations upon treatment with *n*-pentenyl glycosyl donors.<sup>4</sup> Thus, a disarmed donor (NPG<sub>BZ</sub>), e.g. **10**, and/or its orthoester (NPOE) equivalent, e.g. **11**, glycosidated the bold OH overwhelmingly (and frequently exclusively)<sup>5</sup> whereas the armed donor, e.g. **12**, was promiscuous, reacting substantially with both italic and bold-OHs.

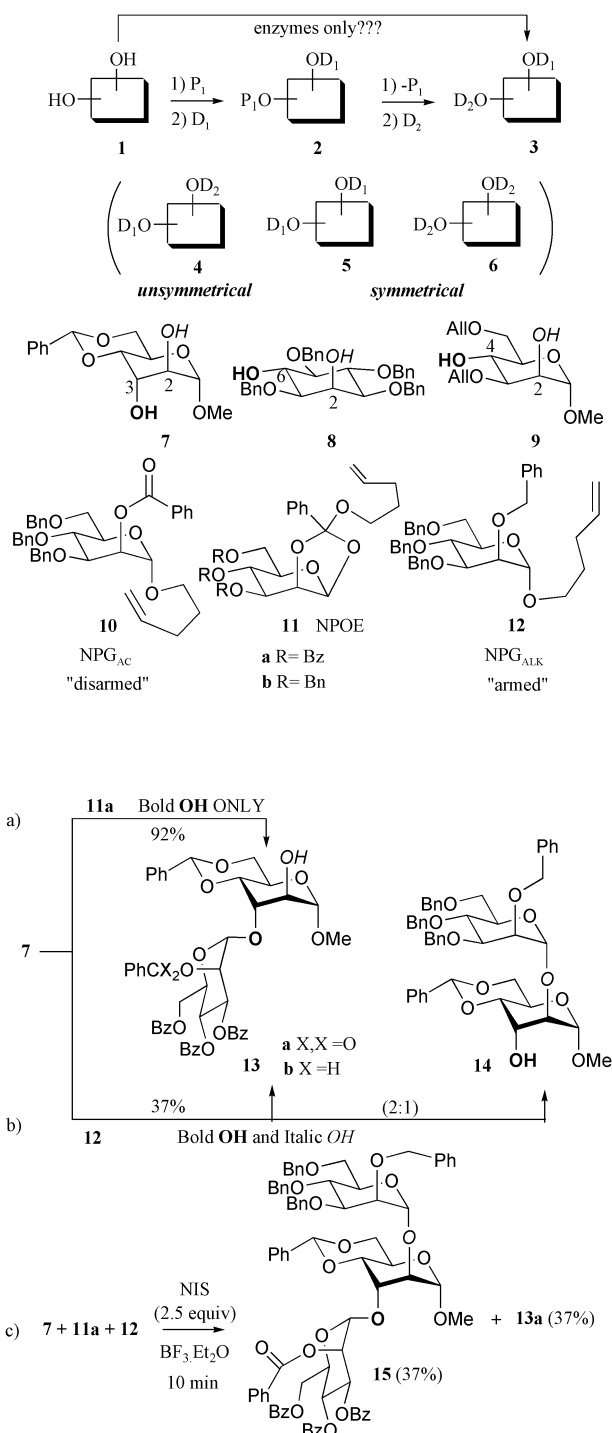
Clearly, these examples indicate that each donor expresses preference for one of the diol –OHs and *vice versa*, the resulting 'match' being evidence for *Reciprocal Donor Acceptor Selectivity* (RDAS).<sup>† 6</sup> A rationalization for these selectivities awaits further insight; but their reality invites immediate practical exploitation.

For in-depth studies, we chose to first examine altroside **7**. This diol had given a 92% yield of **13** as the exclusive product with NPOE **11a**, but a 2 : 1 mixture of **13b** and **14** with the armed NPG<sub>ALK</sub> **12**<sup>4</sup> [Scheme 1 (a and b)]. Notably the preferred site for *both* donors is the bold (C3)-OH which, on the basis of conventional wisdom, did not augur well for differential, double-glycosidation experiments.

Nevertheless, when a 1 : 1.3 : 1.3 mixture of diol **7**, and both donors, (**11a** and **12**) was treated with 2.5 equivalents of NIS and a catalytic amount of BF<sub>3</sub>·Et<sub>2</sub>O for 10 min, trisaccharide **15** and disaccharide **13a**, were obtained as the only products in 37 and 37% yields respectively.

Obviously, the yield of trisaccharide **15**, would be improved if glycosidation of disaccharide **13a** by NPG<sub>ALK</sub> donor **12** could be enhanced. Indeed, entries i–iv in Table 1 show that increasing the concentration of the armed donor **12**, had a salutary effect on the yield of **15**.

An even more interesting set of results arises from our studies on diol **16** (Scheme 2). Differential 3,6-dimannosylation of this mannoside is of interest since the resulting trimannan occurs as



Scheme 1 Glycosidation of altroside **7** with glycosyl donors **11a** and **12**.

**Table 1** One-pot glycosidation of altroside **7** with donors **11a** and **12**.

Entry	<b>7</b> (equiv.)	<b>11a</b> (equiv.)	<b>12</b> (equiv.)	<b>15<sup>a</sup></b> (%)	<b>13a</b> (%)
i	1	1.3	1.3	37	37
ii	1	1	1.6	43	31
iii	1	1	2	52	19
iv	1	1	3	57	16

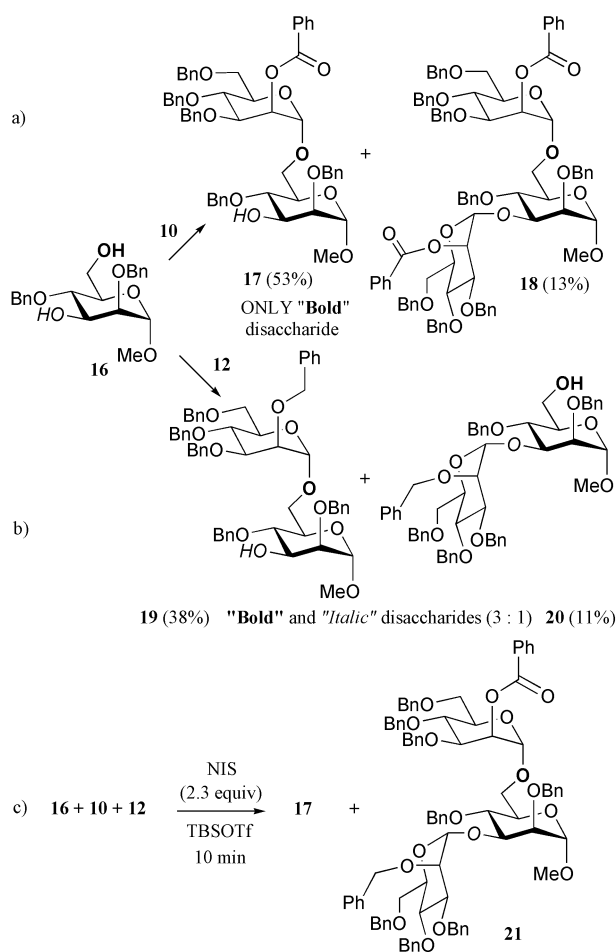
<sup>a</sup> Compound **15** is the *only* trisaccharide (of four possibilities) observed in these optimizations.

a repetitive, interlocking motif in high mannose glycoproteins.<sup>7</sup>

The RDAS preferences of **16** were determined by the previously described equimolar two-component reactions<sup>‡</sup> shown in Scheme 2 (a and b). With the disarmed donor **10**, mannosylation occurred at the bold (C6)-OH only to give **17** in 53% yield, and also the symmetrical trisaccharide **18** in 13% yield—but with *no* evidence (TLC nor NMR) for the dimannan resulting from glycosidation of the italic (C3)-OH. By contrast, the ‘armed’ donor **12** gave a 38% yield of the O6 product, **19**, but also 11% of the O3 regioisomer **20**.

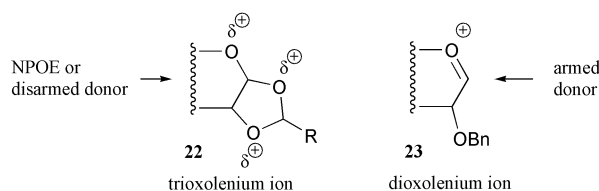
Analysis of the results in Scheme 2 (a and b) according to conventional wisdom, dictates that the preference of both donors, **10** and **12**, for the primary –OH ‘is to be expected’ on the grounds of steric hindrance, and so in contemplating a differential, double-glycosidation experiment, the obvious question was: What will happen when **10** and **12** compete for diol **16**? Our calculations<sup>8</sup> showed that the relative reactivity of these donors ( $k_{12}/k_{10}$ ) is 3.2. Hence, it was expected that O6 mannosylation by the ‘armed’ donor, **12**, would predominate in any trimannan produced.

Surprising disagreement with conventional wisdom is depicted in Scheme 2(c). Thus the 1:1:1 three-component

**Scheme 2** Glycosidation of mannoside **16** with donors **10** and **12**.**Table 2** One-pot glycosidation of mannoside **16** with donors **10** and **12**.

Entry	<b>16</b> (equiv.)	<b>10</b> (equiv.)	<b>12</b> (equiv.)	<b>17<sup>a</sup></b> (%)	<b>21<sup>a</sup></b> (%)
i	1	1	1	36	21
ii	1	1	2	27	25

<sup>a</sup> Compound **22** is the *only* trisaccharide (of four possibilities) observed in these optimizations.

**Scheme 3**

reaction of **10**, **16** and **12** (entry i) gave a single trimannan **22**, in which the *less* reactive donor **10** ended up at O6. But even more surprisingly, the same held true for the single disaccharide, **17**, obtained. As with the altroside study in Scheme 1, an increase in the ratio of the armed donor **12** (entry ii) led to an increase (albeit modest) of trisaccharide **21**—but still *none* of the symmetrical trisaccharide.

In reviewing the above data, we regard it as simply astonishing that even with the audacious disparity in the ratio of donors **12** and **11a** (Table 1, equation iv) or **12** and **10** (Table 2, equation ii), there was absolutely *no* evidence for trisaccharides other than **15** and **21**. In view of the excess of the ‘armed’ donor **12** in these experiments, symmetrical dimannosylation was expected in both Schemes 1 and 2.

A study<sup>8</sup> of the three types of *n*-pentenyl donors used above indicate that their relative reactivities are in the order NPOE > armed > disarmed (e.g. **11** > **12** > **10**). The most and least stable donors therefore give rise to the highly delocalised, more stable intermediate **22**, while the armed donor gives the less stable oxocarbenium ion **23**<sup>9</sup> (Scheme 3). The conclusion from Scheme 1(c) and Scheme 2(c) is that in *competitive* glycosidations, the more stable donor/intermediate (not the most reactive donor) controls regioselectivity, resulting in the formation of the single trisaccharides **15** and **21** and the single disaccharides **13a** and **17**. How the competing OH groups play into this phenomenal regioselectivity awaits clarification.

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## Notes and references

<sup>†</sup> The regioselectivities are the same whether the NPOE or NPG<sub>BZ</sub> is used, although yields and side-products may differ.  
<sup>‡</sup> The structure of the regioisomers was determined, as previously described.<sup>4</sup>

- H. M. I. Osborn and T.H. Khan, *Oligosaccharides. Their synthesis and biological roles*, Oxford University Press, Oxford, 2000, ch. 3.
- H. Paulsen, *Angew. Chem., Int. Ed. Engl.*, 1982, **21**, 152.
- C.-H. Wong, R. L. Halcomb, Y. Ichikawa and T. Kajimoto, *Angew. Chem., Int. Ed. Engl.*, 1995, **34**, 521; S. David, C. Auge and C. Gautheron, *Adv. Carbohydr. Chem. Biochem.*, 1991, **49**, 176.
- B. Fraser-Reid, J. C. Lopez, K. V. Radhakrishnan, M. Mach, U. Schlueter, A. M. Gomez and C. Uriel, *J. Am. Chem. Soc.*, 2002, **124**, 3198.
- G. Anilkumar, L. G. Nair and B. Fraser-Reid, *Org. Lett.*, 2000, **2**, 2587.
- B. Fraser-Reid, J. C. Lopez, K. V. Radhakrishnan, M. Mach, U. Schlueter, A. M. Gomez and C. Uriel, *Can. J. Chem.*, in press.
- J. Montruil, *Adv. Carbohydr. Chem. Biochem.*, 1980, **37**, 157.
- B. G. Wilson and B. Fraser-Reid, *J. Org. Chem.*, 1995, **60**, 317.
- With S. Grimme and M. Piacenza, unpublished results.